

Business Analytics

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Preface

Why this book?

This book is originally intended as material for the BUSI 4720 undergraduate course on Business Analytics. This course is a core, required course for the Bachelor of Commerce program at Memorial University of Newfoundland, Canada. As students receive only a single course in business analytics, and this course is in the fourth and final year of the program, the material coverage is intentionally broad, and covers aspects that may be outside some narrower conceptions of analytics. Additionally, students taking the course generally have little to no exposure to computer applications or statistical software, necessitating a rather comprehensive approach that not only introduces computer and programming basics, such as data and data types that students may encounter in business analytics, but also covers introduction to R and Python as well as a brief coverage of relational and graph databases, that are typically not considered part of business analytics. On the other hand, this course also contains advanced topics, such as interpretable machine learning, analytics at industrial scale, reinforcement and MLOps, that are not usually found in a business analytics course. However, these topics are gaining importance and it is essential that students have at least some exposure to them. In summary, the book was written because no other single book offers the necessary broad perspective.

The book is written from an applied perspective. I believe that students, even business students, should not only be able to talk about analytics, but must also be able to do analytics. This means that, together with the concepts, every chapter also contains R or Python code showing how the concepts can be applied. Looking at this from another perspective, I believe students must not rely solely on software tools and statistical libraries, but it is crucial that they understand, at least in principle and at an intuitive level, how these tools work. This is necessary to allow an informed use of tools, to be able to select the appropriate tool for a given situation, and to be aware of the shortcomings, drawbacks, boundary conditions, and other limitations of tools. In short, formulas in this book are to explain what happens "behind the scenes" of the code, and code is in this book to show how formulas can be applied; both are necessary.

Why these tools?

The focus on R and Python, over commercially available tools, is due to multiple reasons. First, the use of open-source software makes the material more easily accessible to students, independent of the availability of campus-wide licenses, or the use of limited "evaluation" licenses for some commercial tools. A second reason is the cross-platform nature of these software tools. Computing hardware in practice, and in the classroom, is a heterogeneous mix of different chip sets (Intel, Apple/ARM) and different operating systems (Windows, MacOS, Linux, etc.) so that is essential to work with software tools that are available and interoperable across these hardware and operating system platforms. A third reason is that R and Python are widely used in production environments. They tend to be more flexible than commercial offerings, and are also at the forefront of new developments in the area of business analytics. New methods and techniques are typically implemented directly by their inventor in open-source libraries and packages for R or Python, before they mature and are included in commercial offerings. The focus on command line tools is to avoid the complexities of graphical user interfaces that tend to change more rapidly than application programming interfaces (APIs), it is focus on the essentials and not be distracted by graphical environments. Scripting with command line tools generally also leads to better replicability of analyses and easier integration into production environments. For example, while it is all well and good to explore customer purchasing predictions on a small data set using the desktop edition of SPSS (a commercial, graphical, statistics software application), implementing real-time dynamic pricing in the global web-based ordering system will require the model to be implemented and integrated with very different tools.

Instructors: How to use this book?

For instructors, the book is written for a 24 class semester of 75 minutes each (the chapter on visualization should be covered in two classes), with two classes dedicated to mid-term exams. If time is short, some of the later, more advanced chapters could be omitted, for example, the two chapters on reinforcement learning, and/or the chapter on MLOps. A slide set for 22 classes is available, as is a question bank of multiple-choice questions for each chapter, e.g. for quizzes. Each chapter also contains a set of short hands-on exercises that can be used during class to keep students engaged or can form the basis for a computer lab setting. Also available is a set of example exam questions. Given the extensive set of online materials on programming in general, and data science and data analytics in particular, ranging from the traditional <https://stackoverflow.com/> site, to Google and YouTube, to the most recent ChatGPT or other LLMs, it is easy for students to complete any technical homework assignment or course project using such tools. Instructors should therefore focus on data and results interpretation and use new or unpublished data sets, if they wish to set such assessment or evaluation exercises at all. Consequently, the example exam questions are long-answer questions that focus on conceptual understanding of the material, and less on technical programming skills.

Students: How to study?

For students, accompanying this book is a virtual machine with all required software installed and data sets provided. I recommend that students at least run the provided example code to get some "hands-on" with the tools. The best way to learn and understand is to experiment and modify the examples. See what happens when parameters or functions are changed. Ask yourself: Does the result match my expectation? Why or why not? Another way to work with the examples is to make sure you recognize the code elements in the formulas and vice versa. If the formula contains an X , where is this specified in the code or where does it appear in the output? Ensure that you can recognize and make the connection between the conceptual or mathematical level and the implementation in software.

Each chapter contains hands-on exercises. These are relatively simple exercises that build directly on the code presented in a section and require only minor changes or adaptations. These exercises invite experimentation with the code and trying different options. They are highly recommended to further your understanding.

Every chapter also contains a number of review questions. These are there to help check your own understanding. At least read and think about the questions, even if you do not write out any answers.

Many chapters contain pointers to textbooks that formed the basis for the material in this book, and all contain links to online references. These are valuable in that they provide additional, deeper information. And because those resources are written by different authors, they may be easier to understand; at the very least they can provide a different and complementary perspective on the material in this book. Many textbooks that have been used to inform this book are popular or classic textbooks in their own right. Many could form the basis of a somewhat more narrowly conceived course on business analytics. In short, they make for excellent complementary reading and are highly recommended. The vast majority of them are also freely available on the internet.

Additionally, a wealth of information is available in various formats on the internet. This begins with Wikipedia pages, which provide a good introduction to many topics, and material from Wikimedia Commons has been used extensively in this book. Since all the tools used in this book are open-source tools, their web sites provide not only the code, but more importantly, also provide documentation in the form of tutorials, introductions, and detailed programming descriptions. These are all excellent resources. Many researchers and teachers in the area of machine learning have made their materials freely available, for example in their blogs or in YouTube videos and entire YouTube channels. Many of these researchers are active at the forefront of machine learning and are excellent teachers. These resources are valuable resources and provide more depth than offered in this book. At the same time, the current popularity of the topic has also led to some questionable material on the internet, and caution should be exercised when searching for material. Begin your internet search with a trusted source, for example Wikipedia, a well-known researcher, or material from a university instructor active in the field.

What about ChatGPT?

Absolutely consider using your favourite large language model (LLM) for studying this material. For example, ChatGPT is quite good at explaining things. Ask it to explain a code fragment that you copy and paste into it. Ask it to translate R code to Python code or vice versa. Ask it to simplify code for you. Ask it to check code for mistakes. Other ways to engage your favourite LLM are to ask it to quiz you on the material; copy and paste the material into it and then ask it to generate questions and wait for your answer, then to evaluate and correct your answers. Yet another use of an LLM is to ask it to evaluate and correct your answers to the review questions, provide both the question and your answer to the LLM.

These are valuable ways in which you can further your understanding, but keep in mind that LLMs are simply statistical models that predict the most likely next word in the output. As such, they cannot truly reason, they have no intelligence (at least in terms of how we conceive human intelligence) and they make mistakes without being aware of them. So, be careful when you engage them. However, the beauty of using an LLM with computer code is that you can run the code and verify that it does what you expect it to do.

Chapter 1

Introduction

1.1 Introduction

This section describes some of the terminology around the rapidly expanding field of data analytics, business analytics, data science, statistics, machine learning and AI (artificial intelligence).

Data analytics (or simply 'analytics') refers to the broad collection of methods, techniques, and tools to allow humans to make sense of information for purposes of understanding and decision making. Business analytics is the application of data analytics to operational, tactical, or strategic management in businesses and other organizations. Examples are the use of visualization of human resource performance data, trend analysis of outbound logistics costs, prediction of customer demand, fraud analysis of financial transactions, and others.

Data analytics as a broad field is closely related to a range of other fields, such as data management (how best to collect, store, access, and use data in a variety of format), visualization (how best to present data in an easy-to-understand format to generate insights or persuade stakeholders), machine learning (how to train computers to classify data and to make predictions), or text analysis (how to extract meaningful information from natural-language text information). Some argue that these fields are within analytics, but others view them as separate but strongly interrelated. For example, text analysis and machine learning overlap when training computers to predict customer behaviour based on social media post data. Text analysis can provide certain features that characterize or summarize a text, and may be used as input for machine learning. Machine learning models can exploit text-specific features, such as the sentence structure, in making predictions. Training machine learning systems also requires a very large amount of data, so advanced data management techniques are required in order to store and provide this data in an efficient way.

Artificial Intelligence (AI) is a field with a long and varied history, going back to the

1960s. Originally, AI was used for symbolic computations, where researchers attempted to explicitly describe and model human reasoning processes in a computer. In the late 1990s and early 2000s, AI has morphed to focus on statistical models and has most recently become dominated by artificial neural networks (ANN) and deep neural networks (DNN), a field called deep learning. Artificial neural networks, while inspired by the human brain, are essentially statistical models for classification and regression, akin to the simple linear or logistic regression. But whereas the simplest linear regression model may describe a small dataset of hundreds to thousands of observations using just two parameters (the slope and intercept), ANN and DNN are non-linear and highly complex with millions or even billions of parameters and are often trained on billions of observations. However, the main ideas are the same in that the model is trained on or fitted to a data set.

More recently, since about 2020, *generative AI*, that is, AI models and systems that are used to generate text, images, audio or video in response to user input has become synonymous with AI. The rise in popularity of systems such as ChatGPT, Dall-E, and many others, has led to many people equating AI with generative AI.

While *machine learning* may sometimes be viewed synonymously with ANN or DNN, the field of machine learning is broader than just neural network models and concerns the development of methods to make predictions for new observations. Traditional statistical techniques for regression and classification, such as decision trees or support vector machines, are considered part of machine learning, and therefore also part of data analytics. However, these statistical models sometimes take a back seat role compared to ANNs because of the power and flexibility of the latter.

Machine learning and AI are also sometimes viewed synonymously. However, as noted above, there are subfields of AI that are not concerned with machine learning and prediction. First, research into symbolic reasoning is still ongoing. Second, methods such as reinforcement learning, which focuses not on making predictions, but on prescribing optimal courses of action, are considered part of machine learning and AI.

Big Data was an important topic in the early 2000s and 2010s but is now often considered a sub-field of data analytics or data science. The motivation for Big Data was the recognition that the volume of data produced and available for analysis has been growing exponentially since the 1990s. This has spurred the development of advanced data management methods, techniques, and tools, such as distributed file systems and databases. The velocity of data, that is, its rate of production, has also increased greatly since the 1990s. Processing the data often has to occur in real-time, leading to development of techniques and tools that can analyze data "on-the-fly" ("stream processing").

Finally, the term *data science* is often used in a less applied and more scientific or research and developmental way than the term data analytics, to characterize the development, rather than application, of methods, techniques, and tools.

1.2 Methods, Techniques, and Tools

In the context of data science and data analytics, methods, techniques, and tools play distinct roles in the process of extracting insights from data. *Methods* encompass overarching approaches and strategies, providing a systematic framework for tasks such as data exploration, modeling, and analysis. These high-level methodologies guide data scientists in formulating a structured plan for addressing specific challenges or achieving analytical goals.

Techniques in data science are the specific procedures and practices employed within the broader methodological framework. These practical and detailed approaches are applied to handle particular aspects of the data analysis process. For instance, in exploratory data analysis, techniques like histograms or scatter plots are used to visually inspect data distributions or relationships between variables and in classification, a naive Bayes classifier is considered a specific technique.

Tools in data science refer to the instruments and software applications that facilitate the practical application of methods and techniques. These can range from programming languages like Python and R, statistical packages such as Pandas and SciPy, to visualization tools like Shiny or Matplotlib. The selection of appropriate tools is crucial for efficiently executing data science tasks and optimizing the workflow.

1.3 Types of Analytics

There are different "types" of analytics that have different aims.

Descriptive Analytics describes "what is". It typically provides summaries of the data, makes comparisons between different types of entities or measurements, may identify historical trends, or provide rankings of observations or measurements. An example is the identification and comparison of current and historical costs to manufacture different widgets in different plants. Another example are the identification of the top-grossing sales people in the organization for specific product types, or calculating the mean cycle time of the order-to-cash business process for different time periods. Descriptive analytics is therefore important in a business context and organizations expend a great deal of money, time, and effort on technologies such as report generating tools to support this type of analytics.

Predictive Analytics describes "what may be" in the future. It typically builds a *model* based on past data to predict future cases/events/outcomes. As a simple example, consider a linear regression model that predicts the overall spending of a customer from their income. One can build a linear model with parameters for the intercept and slope and then train the model on customers whose spending and income are known. Training means to determine the two parameters of the model so that they model best fits the data. Given a trained model, one can then predict the overall spending of a new customer given their income. Of course, the models can be much more complex than a

simple linear regression, and often have tens, hundreds, thousands, or even millions of parameters, but the principle of model-based predictive analytics remains the same.

Prescriptive Analytics describes "what should be done". Similar to predictive analytics, a model is usually built from past data. However, the model must now also consider which actions can be taken (and were taken for the past training data). In reinforcement learning, a popular prescriptive analytics technique, one assumes that an *agent* can observe the state of an *environment*, can take *actions* based on the observed state, and receives a (positive or negative) reward from the environment after taking an action. Actions may change the state of the environment. The agent must learn to identify those actions in each state that give it the maximum rewards. The optimal actions are called a *policy* and can then be used to prescribe which action to take in which state.

Visual Analytics describes the use of graphs (plots, diagrams) to visualize information for exploring data and gaining insight. Visual analytics makes use of the human ability to visually identify trends, make comparisons, etc. In effect, this type of analytics supports humans in their tasks. It employs no mathematical or statistical models. However, humans may identify features of the data such as trends, comparisons, etc. with visual analytics that may then be tested with statistical models or form the basis on which to build predictive or prescriptive analytics models.

1.4 Machine Learning

In the context of analytics, machine learning (sometimes simply called "learning") can be divided into supervised and unsupervised learning. *Supervised learning* is typically based on a parameterized statistical or mathematical model. For example, a simple linear regression is based on two parameters, the slope and intercept. Models are trained (that is, parameters are estimated) by adjusting the model parameters so that the model's output (e.g. the predicted value " \hat{y} " in a linear regression) for each given input (e.g. the " x " value in a linear regression) matches the actually observed outcome (the " y " value in a linear regression). Supervised learning assumes that a correct/observed outcome is available for all inputs. Linear regression is a very simple supervised learning approach with a simple statistical model; on the other extreme there are generative pre-trained transformer (GPT) models that predict words for generating text, and which contain billions of parameters and non-linear relationships among them.

Unsupervised learning on the other hand does not require outcome values and only requires "input" (" x ") values. Typical unsupervised learning tasks are clustering and dimensionality reduction. For example, one may form clusters of widgets at the end of a production line based on how similar the widgets are on different characteristics that were measured by sensors. These clusters could then, for example, be interpreted as quality grades. Similarly, customers may be clustered based on their past transactions or purchases. Different marketing strategies may then be applied to each cluster. The aim of dimensionality reduction is to be able to describe a data set with many variables

by using only a few variables. Imagine that having hundreds of different characteristics of widgets. One might then wish to simplify and identify fewer, perhaps as few as two or three, combinations of the original characteristics that provide the same information about the widgets.

1.5 Analytics is not Statistics

Analytics and statistics use the same kinds of mathematical models. However, "traditional" *statistics* focuses on sample and population characteristics, such as means, slopes, intercepts, and others, of a sample or a population that are represented as parameters of a mathematical model. Statistics aims to identify and explain the data generating mechanism, i.e. the "real world" or population. In particular, *inferential statistics* is used to generalize from a sample to a population. Importantly, statistics is typically not concerned with individual cases or individual observations.

In contrast, data analytics, especially predictive analytics, focuses on predicting the outcome of an individual case or observation. Analytics is pragmatic, in that models are considered useful tools and do not need to faithfully describe the "real world" or the data generating mechanism, as long as they make good predictions or are otherwise useful for their purpose. Consequently, there is no inference from sample to population, because the model does not claim to describe a population. The quality of a model is determined not by its fit with the observed data, but by its precision or accuracy when predicting specific observations.

1.6 Tools used in this Course

The software used in this course is open-source and free software. Open-source software (OSS) embodies a collaborative approach to software development, allowing users to access, modify, and distribute the source code freely¹. This approach promotes transparency, enabling users to inspect the code, modify, adapt, fix and extend it, and contribute to its improvement.

Free software, as defined by the Free Software Foundation (FSF), goes beyond accessibility, emphasizing users' fundamental freedoms to run, study, modify, and share the software². Contrary to the common misconception, "free" in this context pertains to freedom, not necessarily zero cost. The ethical philosophy behind free software underscores the importance of user control over technology.

The term FOSS, or Free and Open-Source Software³, serves as an inclusive label encompassing both the principles of free software and the collaborative nature of open-source software. FOSS encourages a shared approach to software development, emphasizing not only the technical benefits of open code but also the ethical imperative of user freedom and community-driven innovation.

¹https://en.wikipedia.org/wiki/Open-source_software

²https://en.wikipedia.org/wiki/Free_software

³https://en.wikipedia.org/wiki/Free_and_open-source_software

Free and Open-Source Software (FOSS) *licenses* are legal agreements that govern the use, modification, and distribution of open-source software. These licenses play a crucial role in preserving the core principles of freedom, transparency, and collaboration within the open-source community. Common characteristics of FOSS licenses are:

Freedom to Use FOSS licenses grant users the freedom to use the software for any purpose without any restrictions.

Freedom to Study Users have the right to access and study the source code of the software. This transparency allows for a deeper understanding of how the software functions.

Freedom to Modify FOSS licenses typically allow users to modify the source code according to their needs. This encourages innovation, customization, and adaptation of the software.

Freedom to Share Users can distribute both the original and modified versions of the software, fostering a collaborative environment. This freedom to share is fundamental to the open-source philosophy.

Copyleft Licenses Some FOSS licenses, such as the GNU General Public License (GPL), include copyleft provisions. Copyleft ensures that any derivative works or modifications are also subject to the same open-source terms. This prevents the software from being incorporated into proprietary projects without maintaining open-source characteristics.

Permissive Licenses On the other hand, permissive licenses, like the MIT License and the Apache License, allow for more flexibility. They permit the use of the software in proprietary projects without imposing the requirement to open-source the derived code.

While a lot of free and open-source software used to be developed by individuals, increasingly FOSS is developed by companies whose business model rests either on providing paid support for such tools, or on providing paid hosted versions of the software, or on providing non-free extensions for their FOSS software. In other cases, companies provide software developer time to important projects that benefit themselves.

The R System



R is a programming language and free software environment designed for statistical computing and graphics.

The history of R begins in the early 1990s at the University of Auckland, New Zealand. Ross Ihaka and Robert Gentleman, two statisticians, set out to create a programming language that would

R	4.1.2	www.r-project.org
dplyr	1.1.3	www.tidyverse.org
tidyverse	1.3.0	www.tidyverse.org
ggplot2	3.4.4	www.tidyverse.org
Python	3.8	www.python.org
numpy	1.24.4	numpy.org
pandas	2.0.3	pandas.pydata.org
plotly	5.18.0	plotly.com
tensorflow	2.13.1	www.tensorflow.org
Postgres	16.0-1	www.postgresql.org
pgAdmin4	7.8	www.pgadmin.org
PyCharm	2023.2.3	www.jetbrains.com/pycharm/
Jupyterlab	4.0.7-1	//github.com/jupyterlab/jupyterlab-desktop
Neo4J	5.14.0	www.neo4j.com

Table 1.1: Software used in this course

make data analysis and visualization more accessible. They released the first version of R in 1995, and it quickly gained traction within the academic community.

Over the years, R evolved and expanded its capabilities, thanks to the collaborative efforts of statisticians, data scientists, and programmers worldwide. The Comprehensive R Archive Network (CRAN) was established to serve as a hub for R packages, fostering a community-driven approach to software development.

The open-source nature of R played a pivotal role in its success. As more people embraced it, R became not just a statistical tool but a versatile platform for data analysis, machine learning, and graphical exploration. Its popularity soared in both academic and industry settings, with businesses recognizing its potential for extracting meaningful insights from data.

R continues to thrive as a dynamic and evolving tool in the world of data science. Its rich ecosystem of packages and active community ensure that it remains at the forefront of statistical computing and analysis.

One of the key hubs of the R community is the Comprehensive R Archive Network (CRAN), where thousands of R packages are hosted. These packages cover a vast array of topics, from basic statistical functions to cutting-edge machine learning algorithms. The "open-source" ethos is strong here, with contributors from around the globe actively developing and maintaining packages.

Stack Overflow⁴ and other online forums serve as virtual interchanges of ideas where R users exchange knowledge and troubleshoot problems.

The tidyverse⁵ is a comprehensive collection of R packages that share a common philosophy and syntax, designed to streamline and enhance the data analysis workflow.

⁴<https://stackoverflow.com/collectives/r-language>

⁵<https://www.tidyverse.org/>

Developed by Hadley Wickham and his collaborators, the tidyverse promotes a principled approach to data manipulation, visualization, and exploration. Key components are:

ggplot2 A sophisticated and flexible plotting system, ggplot2 enables the creation of intricate and publication-ready visualizations. Its grammar of graphics approach provides a consistent framework for constructing a wide range of plots.

dplyr This package serves as a cornerstone for data manipulation, offering a set of succinct and expressive verbs for tasks such as filtering, grouping, summarizing, and joining datasets. Its syntax facilitates a more intuitive and readable coding style.

tidyverse Complementing dplyr, tidyverse focuses on reshaping and tidying data. It provides functions like ‘gather()’ and ‘spread()’ to efficiently restructure datasets, ensuring they adhere to the principles of tidy data.

readr A fast and user-friendly package for reading and parsing data from various file formats. readr’s emphasis on speed and consistency makes it a reliable choice for importing datasets seamlessly.

The tidyverse’s cohesive design and interoperability between packages make it a popular choice for data scientists and analysts seeking an efficient and coherent ecosystem for their R-based projects.

Python

Python⁶, conceived by Guido van Rossum in the late 1980s and released in 1991, has evolved into a versatile and influential programming language. Known for its readability and clean syntax, Python prioritizes simplicity and ease of use, making it accessible to both beginners and seasoned developers. The Python Software Foundation now oversees its development, ensuring that it remains free, open, and continually improved by a global network of contributors.



One of Python’s key strengths is its versatility. It serves as a general-purpose language, excelling in web development, data analysis, artificial intelligence, scientific computing, and more. Its extensive standard library and a rich ecosystem of third-party packages contribute to its adaptability across diverse domains.

Python’s readability, enforced by the use of indentation for block delimiters, facilitates code comprehension and maintenance. This, coupled with a strong emphasis on code readability and maintainability, has contributed to its popularity among developers.

⁶www.python.org

Python's adoption in data science has surged, with libraries such as NumPy, pandas, and scikit-learn forming the backbone of numerous data analytics and machine learning projects.

The language's cross-platform compatibility, supported by its interpreted nature, allows developers to write code once and run it on various operating systems without modification. This, combined with a vast and active community, ensures that Python remains at the forefront of technological advancements.

In the Python ecosystem, packages play a crucial role in extending the language's functionality and addressing specific programming needs. Python packages are collections of modules, scripts, and other resources that facilitate the development of reusable and modular code. Pip (Package Installer for Python) is the default package installer for Python, allowing users to easily install, upgrade, and manage Python packages. It simplifies the process of fetching and installing packages from the Python Package Index (PyPI) and other repositories. PyPI is the official repository for Python packages, hosting a vast collection of open-source Python software.

NumPy NumPy, short for Numerical Python, is a fundamental library in the Python ecosystem for numerical computing. Developed to facilitate array operations, mathematical functions, and linear algebra capabilities, NumPy provides a foundation for scientific and data-intensive applications.

Launched in 2005 by Travis Olliphant, NumPy has become a cornerstone in the Python data science stack. Its core feature is the ndarray, a multidimensional array object that enables efficient manipulation of large datasets. NumPy's array-oriented computing paradigm enhances performance and readability, making it a preferred choice for numerical tasks.

Pandas Pandas, a Python library introduced in 2008 by Wes McKinney, is an important tool in data manipulation and analysis. Designed to provide high-performance, easy-to-use data structures, Pandas simplifies the handling of structured data and time series.

At its core are two primary data structures: the Series, a one-dimensional labeled array, and the DataFrame, a two-dimensional table with labeled axes. These structures, built on top of NumPy arrays, empower users to perform a range of operations from basic data cleaning to complex analytics, with a concise and expressive syntax.

Pandas can handle missing data gracefully and offers easy to use tools for reshaping, grouping, and aggregating data. Its integration with other Python libraries, coupled with efficient indexing and alignment features, makes it a good choice for data scientists, analysts, and researchers working with heterogeneous and large datasets.

Tensorflow and Keras TensorFlow is an open-source machine learning framework initially developed by the Google Brain team. It provides a comprehensive set of



tools and libraries for building and deploying machine learning models. TensorFlow facilitates the creation of artificial neural networks and other machine learning models through a flexible and scalable platform.

Keras is an open-source high-level neural networks API written in Python. Originally developed as an independent library, it has become an integral part of TensorFlow, serving as its official high-level programming interface. Keras simplifies the process of building, training, and deploying neural networks, making it accessible to developers of different skill levels. While Keras is designed to be user-friendly and concise, TensorFlow provides a more extensive and low-level framework for those requiring greater flexibility in model design and customization.

PyCharm PyCharm by JetBrains is an integrated development environment (IDE) for Python developers. PyCharm Community is its free and open-source version. The IDE includes a code editor with syntax highlighting, code completion, and error checking, supporting developers in writing clean and efficient code. PyCharm's code navigation and refactoring tools facilitate easy exploration and improvement of code bases. The built-in debugger and seamless integration with testing frameworks like pytest enhance debugging and testing capabilities. PyCharm also supports version control systems, including Git and Mercurial, promoting collaborative development. PyCharm has a user-friendly interface, continuous updates, and active support.



Jupyter, JupyterLab, JupyterLab Desktop Jupyter Notebooks are interactive computing environments that allow users to create and share documents containing live code, equations, visualizations, and narrative text. Originally developed for Python, Jupyter Notebooks support multiple programming languages through various kernels. The notebooks are structured as a series of cells, where each cell can contain code, markdown text, or rich media elements. Users can execute code cells interactively, see immediate outputs, and create a seamless blend of code and documentation.



JupyterLab is an extensible web-based interactive computing environment developed by the Jupyter Project. It serves as the next-generation interface for Jupyter Notebooks, offering a more versatile and powerful environment. JupyterLab provides a flexible and modular interface where users can arrange documents, notebooks, terminals, and custom components in a tabbed layout. It supports multiple Jupyter Notebooks simultaneously and allows for drag-and-drop functionality to rearrange and organize the workspace. JupyterLab's extensibility comes from its plugin system, enabling users to add new features and customize the environment to suit their workflows.

JupyterLab Desktop refers to the stand alone application version of JupyterLab that runs on a user's desktop rather than in a web browser. It offers the same rich features

as the web-based version but provides a stand alone application that can be launched independently. JupyterLab Desktop enhances user accessibility and convenience, providing a familiar desktop application experience for working with Jupyter Notebooks and other interactive computing tasks.

PostgreSQL



PostgreSQL is an open-source relational database management system (RDBMS) whose history goes back to the mid-1980s when a team led by Michael Stonebraker at the University of California, Berkeley laid the foundation for what would later become PostgreSQL.

PostgreSQL has been characterized by standards compliance, extensibility, and robustness. Over the years, it has evolved into a feature-rich and reliable database system that caters to a wide range of applications. One of PostgreSQL's key strengths lies in its extensibility and support for custom data types, operators, and functions. This flexibility empowers developers to model and store data in ways that suit the specific needs of their applications. Furthermore, PostgreSQL boasts support for advanced indexing techniques, complex queries, and transactional consistency, making it well-suited for high-performance and mission-critical environments.

The commitment to data integrity is a hallmark of PostgreSQL. It provides support for ACID (Atomicity, Consistency, Isolation, Durability) properties, ensuring the reliability of transactions. Additionally, features like point-in-time recovery and built-in replication mechanisms contribute to the system's resilience and availability.

Neo4J



Neo4j is a graph database management system, designed to handle and store data in a graph structure rather than in traditional tables. In Neo4j, data is represented as nodes, edges, and properties. Nodes typically represent entities such as people, businesses, accounts, or any

other item you might find in a dataset. Relationships provide the connections between these nodes, akin to how foreign keys work in relational databases, but with a more natural and direct approach to represent how items are related. This structure makes it particularly efficient for querying complex and deeply interconnected data.

One of the key strengths of Neo4j is its powerful query language, Cypher, which allows for expressive and efficient querying and manipulation of the graph data. Cypher is designed to be intuitive and readable, focusing on the clarity of expressing what data to retrieve or how to manipulate the data, rather than how to navigate the structure. This makes it easier to model complex relationships and query them efficiently. Neo4j is a popular choice for applications that require complex data relationships and network analysis, such as social networks, recommendation systems, and fraud detection.

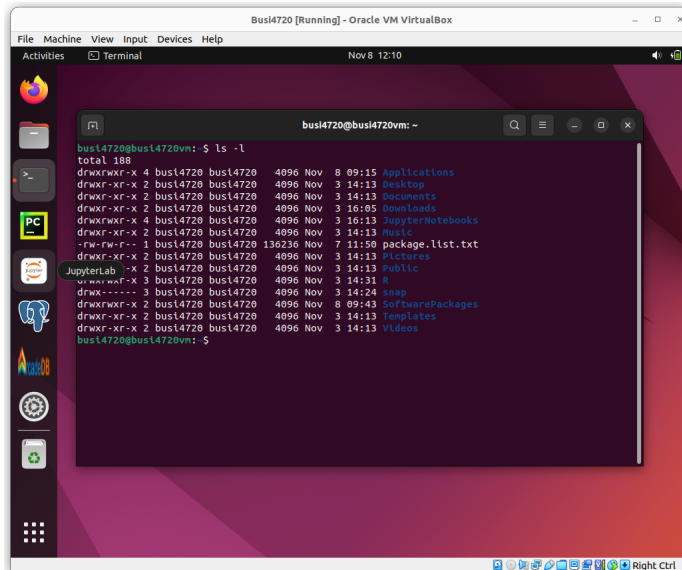
1.7 Ubuntu Linux



Ubuntu Linux, developed and distributed by Canonical Ltd., has emerged as a popular distribution within the Linux ecosystem since its first release in 2004. Through its commitment to ease-of-use and pragmatic approach to hardware compatibility, Ubuntu has played a significant role in popularizing Linux as a viable alternative to proprietary operating systems. The operating system's development is steered by a global community of contributors who collaborate on various aspects, from bug fixes to the introduction of new features.

One distinguishing feature of Ubuntu is its emphasis on regular and predictable release cycles. This approach ensures that users have access to the latest software updates, security patches, and improvements. Ubuntu's Long Term Support (LTS) releases, occurring every two years, provide a support window of 5 years, offering stability and reliability for enterprises and users seeking a more predictable environment.

The desktop edition of Ubuntu employs the GNOME desktop environment, providing an intuitive and user-friendly interface. Ubuntu's package management system, APT, simplifies the installation and removal of software packages. It uses the "apt" command line tool or the "Synaptic" graphical interface to APT. The extensive and well-maintained software repositories contribute to Ubuntu's versatility, allowing users to access a rich ecosystem of applications without the need for extensive manual configurations. Because Ubuntu is based on Debian, it can use Debian packages and packages in the Debian package format.



As a multi-user operating system, Ubuntu provides support for file permissions for each

user, user home directories, and user privileges. User home directories are typically located in `/home/<userName>/` and the `sudo` ("superuser do") command may be used to execute commands as super user (equivalent to the "root" user in Linux terminology).

1.8 Virtual Machines

Virtualization with virtual machines allows users to create and run virtual computers on their own computers, enabling the installation and operation of multiple operating systems simultaneously. The "real" system that is running the virtualization software application is called the host system or host operating system, while the virtual computer running inside the virtualization software is called the guest system, or guest operating system. In effect, the virtualization software pretends to be an actual computer to the guest system, and the guest system is a complete operating system, such as Windows or Linux.



VirtualBox is a free and open-source virtualization software application developed by Oracle. VirtualBox is available for host systems with an Intel or AMD processor, running Windows, MacOS, or Linux operating systems. VirtualBox supports a wide range of guest operating systems, including various versions of Windows, Linux distributions, MacOS, BSD, and others. VirtualBox provides "Guest Additions," which are additional software packages that can be installed on the guest operating system. These additions enhance the performance and integration between the host and guest systems, providing features like seamless mouse integration, shared folders, and improved graphics support.

VMWare Fusion is a proprietary virtualization software owned by Broadcom. VMWare Fusion is available for Apple Mac computers running either Intel processors (before circa 2021) or the later Apple M1, M2 or M3 processors (after circa 2021). Similar to VirtualBox, it allows users to create and run virtual machines on their computer and provides a way to share folders from the host system to the guest system, and to copy and paste from host to guest and vice versa.

Virtual machine files for use with VirtualBox and VMWare Fusion are provided^a that contains an Ubuntu system with all required software installed. If you wish to use this, you must install VirtualBox or VMWare Fusion on your computer, then download the corresponding virtual machine file and import it into the virtualization software application.

The username is **busi4720** and the password is **busi4720**. Whenever a password is required, you should enter **busi4720**.

^a<https://evermann.ca/busi4720.html>

If you do NOT wish to use the VirtualBox Appliance, you should download and install all software to your computer from the sources indicated in Table 1.1 in (at least) the versions indicated in the table.

1.9 The Ubuntu Command Line (also for Mac Users)

This tutorial provides a very brief introduction to the Ubuntu command line ("terminal"). The command line, also called a "shell" is by default the "bash" shell (Bourne-again shell; a pun on the earlier Bourne shell). In Ubuntu, you can open the Terminal application using the key combination **Ctrl-Alt-T**, or by selecting the Terminal application icon from the side bar or the application list. You can also open a Terminal from the file browser.

Note: The default shell in the MacOS terminal is the "zsh" and behaves slightly differently from the bash shell. You can work with a "bash" shell by typing the `bash` command in a MacOS terminal.

Bash will show you a *command prompt* that indicates your username ("busi4720"), the name of the computer ("busi4720vm") and your current working directory ("~") followed by the dollar sign "\$".

Print the working directory by typing the `pwd` command and then pressing the **Return** or **Enter** key:

```
busi4720@busi4720vm:~$ pwd
/home/busi4720
```

Make a folder/directory with the `mkdir` command (in your current working directory):

```
busi4720@busi4720vm:~$ mkdir someFolder
```

Change the working directory to the folder you have just created with the `cd` command. Note how the Bash command prompt indicates your new working directory.

```
busi4720@busi4720vm:~$ cd someFolder
busi4720@busi4720vm:~/someFolder$ cd ..
busi4720@busi4720vm:~$ cd ~
```

The following special characters can be used when specifying folders/directories and paths:

~	User home directory
.	Current directory
..	Upwards in the directory tree
/	Root of directory tree

Here are some tips that make working with the shell a lot easier:

- Autocompletion of file names is available with the “**Tab**” key. When multiple file names exist that match what you have entered so far, you can enter further characters of a file name to disambiguate and press the **Tab** key again for further autocompletion.
- You can recall earlier commands with the “**Up Arrow**” key. By default, the shell stores the last 1000 commands.
- You can search earlier commands with the “**Ctrl-R**” key. You are then prompted to search by typing in characters to find commands. The shell finds the most recent command that contains the characters you entered. At any time you can press **Ctrl-R** again to find earlier matches to your command search.
- Because the usual keys **Ctrl-X**, **Ctrl-C**, **Ctrl-V** for cutting, copying, and pasting text have different functions in the shell, you can cut, copy, and paste with the **Ctrl-Shift-X**, **Ctrl-Shift-C**, and **Ctrl-Shift-V** keys.

List folder/directory contents using the **ls** command. The option **-l** for the command indicates that you would like to see long results.

```
busi4720@busi4720vm:~$ ls -l ~/Applications
total 8
drwxrwxr-x 7 busi4720 busi4720 4096 Nov  8 12:05 arcadedb-23.10.1
drwxr-xr-x 8 busi4720 busi4720 4096 Nov  7 11:45 pycharm-community-2
```

The results show the total size in kB, and a list of entries:

- Type of entry (“d” = directory)
- Permissions for owner of the file (“rwx”), users in the same user group as the owner (“r-x”) and other users (“r-x”): r indicates read access, w indicates write access, x indicates permission to run the application or enter/view a directory (with **cd** or **ls**), and a – indicates the lack of the corresponding permission.
- Names of owner and groups (“busi4720”)
- Size (in bytes)
- Last modification date and time
- File or directory name

Print a string of text using the **echo** command:

```
$ echo "To be or not to be"
To be or not to be
```

Redirect the output of the `echo` command to a file using the *redirect* symbol "`>`". You can redirect the output of any command this way. Use `»` to redirect and append, instead of overwriting a file.

```
$ echo "To be or not to be" > someFile.txt
$ ls -l someFile.txt
-rw-rw-r-- 1 busi4720 busi4720 19 Nov  8 14:50 someFile.txt
```

Print contents of a file ("concatenate") using the `cat` command. You can concatenate multiple files by specifying them all (this is why the command is called "concatenate"):

```
$ cat someFile.txt
To be or not to be
```

If you would like to see the contents of a file page by page or line by line, use the `less` command (a pun on "less is more" and the earlier "more" command that did the same). You will be shown the contents and can navigate up and down with the usual arrow keys.

```
$ less someFile.txt
```

Copy a file using the `cp` command:

```
$ cp someFile.txt someCopy.txt
```

Move a file to a new location (folder/directory) using the `mv` command:

```
$ mv someCopy.txt ~/someFolder
```

Renaming is moving. When you want to rename a file, move it to a new file name:

```
$ mv someFile.txt newName.txt
```

Remove (delete) a file using the `rm` command:

```
$ rm someFolder/someFile.txt
```

Remove a directory recursively (i.e. remove all its contents first):

```
$ rm -r ~/someFolder
```

Use this very carefully! You could inadvertently delete all your files. The shell will delete files and folders immediately and irrevocably. There is no "undoing" this.

View the command line history with the `history` command. Remember that you can redirect this output to a file if you wish or use a pipe to pipe it into the less command (see below).

```
$ history
1 echo "To be or not to be"
2 echo "To be or not to be" > someFile.txt
3 ls -l someFile.txt
4 less someFile.txt
5 cat someFile.txt
...
```

Management of file permissions is done using the `chmod` command. You can grant and revoke read, write, and execute permissions for yourself, your group members, and other users. Remove write permission for yourself by using the `-w` option and specifying the filename:

```
$ chmod -w newName.txt
```

Add write permissions using the `+w` option:

```
$ chmod +w newName.txt
```

Add execute permissions using the `+x` option:

```
$ chmod +x newName.txt
```

If you are stuck on how to use a command or wish to see all its options and capabilities, you can get the manual for a command using the `man` command:

```
$ man ls
```

If you can't quite remember which command to use, you can search for commands using keywords with the `apropos` command:

```
busi4720@busi4720vm:~$ apropos python
keyring (1)           - Python-Keyring command-line utility
pdb3 (1)              - the Python debugger
pdb3.10 (1)            - the Python debugger
pip (1)                - A tool for installing and managing Python p...
pip3 (1)               - A tool for installing and managing Python p...
py3compile (1)          - byte compile Python 3 source files
py3versions (1)         - print python3 version information
pydoc3 (1)              - the Python documentation tool
pydoc3.10 (1)            - the Python documentation tool
pygettext3 (1)            - Python equivalent of xgettext(1)
pygettext3.10 (1)          - Python equivalent of xgettext(1)
python (1)                - an interpreted, interactive, object-oriente...
python3.10-config (1)      - output build options for python C/C++ exte...
python3 (1)               - an interpreted, interactive, object-oriente...
...
...
```

You can see a list of all processes currently running using the `ps` command. The results show the process identifier (PID), the console from which you started the process, the computing time it has consumed, and the command that was used to start the process. Add the `a` and `x` option to see *all* the processes running on your computer, not just the processes you have started.

```
$ ps ax
  PID TTY          TIME CMD
 3024 pts/0    00:00:00 bash
 3151 pts/0    00:00:00 ps
```

The `grep` command is useful to find something in a file or input stream. Use it as in the following example in a pipe:

```
$ cat newName.txt | grep be
$ ls -l | grep .txt
$ history | grep .txt
```

Note: The vertical bar is called a "*pipe*", it pipes the output of one command as input into the next one

The following are further beginner-level tutorials on using the command line on Ubuntu (or really any Linux distribution):

- [Ubuntu command line for beginners](#)
- [Linux command line primer](#)
- [Getting started with Linux](#)

1.10 Review Questions

General

1. What is the definition of "data analytics"? What is "business analytics"?
2. What is the relationship between data management and analytics?
3. Give examples of areas related to analytics and their relationships.
4. Why is text analysis mentioned in connection with machine learning?
5. What are artificial neural networks (ANN) and deep neural networks (DNN), and how are they the same and how are they different from linear regression models?
6. In what way does AI include areas beyond machine learning?
7. Characterize Big Data and its focus.
8. Provide an example that illustrates areas of AI outside of data analytics.
9. Define techniques in the context of data science.
10. Provide an example of a technique used in exploratory data analysis.
11. Mention a few examples of tools used in data science.
12. Explain the relationship between methods and techniques in data science.
13. Provide examples of tasks or challenges that high-level methodologies (methods) might address in data science.
14. Summarize the roles of methods, techniques, and tools in the context of data science.

Types of Analytics

15. What is the primary purpose of descriptive analytics? Give an example of how it might be used in a business context?
16. Explain how predictive analytics differs from descriptive analytics. Illustrate with an example how a simple linear regression model can be used in predictive analytics.
17. Describe prescriptive analytics and how it differs from predictive analytics.
18. What is visual analytics and how does it support human tasks in data analysis? Discuss how it can contribute to other types of analytics.
19. Compare and contrast predictive and prescriptive analytics in terms of their approach and end goals. How do they both utilize past data?
20. If a company wants to understand its sales performance over the last five years, which type of analytics would be most appropriate and why?
21. Imagine a scenario where a company needs to decide on future marketing strategies. Which type of analytics would be most beneficial for them and how might it be implemented?

Learning

22. What is supervised learning in machine learning, and how does it differ from unsupervised learning?
23. Describe how a simple linear regression model works in supervised learning. What are the key parameters in this model?

24. Compare the simplicity of a linear regression model with the complexity of a model like GPT in supervised learning. What are the key differences?
25. Provide examples of tasks that are typically performed using unsupervised learning.
26. How is clustering used in unsupervised learning, and can you give an example of its application in a business context?
27. Explain the concept of dimensionality reduction in unsupervised learning and its potential benefits.

1.11 Hands-On Exercises

The following are a set of connected exercises to help you practice your command line skills. Do them in the order listed.

1. Navigation and Listing
 - (a) Open the terminal and use the `pwd` command to print the current working directory.
 - (b) Use `ls` to list the contents of the current directory.
 - (c) Create a new directory named "Exercise1" using `mkdir`.
 - (d) Navigate into the "Exercise1" directory using `cd`.
2. File Manipulation
 - (e) Create a new file named "file1.txt" inside the "Exercise1" directory using `touch`.
 - (f) Use `cat` to display the contents of "file1.txt".
 - (g) Append the text "Hello, Bash!" to "file1.txt" using `echo` and `>>`.
 - (h) Display the updated contents of "file1.txt" using `cat`.
3. Removing and Renaming
 - (i) Remove "file1.txt" using the `rm` command.
 - (j) Create a copy of the "Exercise1" directory named "Exercise1_backup" using `cp -r`.
 - (k) Remove the original "Exercise1" directory using `rm -r`.
4. Directory Manipulation
 - (l) Recreate the "Exercise1" directory.
 - (m) Create three subdirectories inside "Exercise1" named "Subdir1", "Subdir2", and "Subdir3" using `mkdir`.
 - (n) List the contents of "Exercise1" to verify the creation of subdirectories.
5. Searching and Filtering
 - (o) Create a file named "keywords.txt" inside "Exercise1" and add some random text.
 - (p) Use `grep` to search for a specific word (e.g., "Bash") in "keywords.txt".

- (q) Create a new file named "filtered.txt" and use `grep` to filter lines containing the word you searched for in "keywords.txt".

6. Process Management

- (r) Use `ps` to display information about the current processes running on your system.
- (s) Use `ps aux | grep bash` to filter and display information about Bash processes.

7. Cleanup

- (t) Remove the entire "Exercise1" directory and its contents using `rm -r`.
- (u) Confirm that the "Exercise1" directory no longer exists by listing the contents of the current directory.

Chapter 2

Data, Data Types, Data Quality

2.1 Introduction

Business analytics is the use of data for understanding, description, prediction, prescription and decision making. Hence, it is important to understand the different types of data and the variety of formats in which they can exist.

This section first introduces primitive data types, such as numbers and text. There are many complexities to be aware of that can make analytics challenging. Next, complex data types are introduced, such as tables, documents, and graphs, which are useful in describing complex information, such as customer purchase history in tables, product descriptions in documents, or supply chain logistics in a graph. Then, unstructured data in the form of text, images, and audio/video information is explained. Data in these formats may be market information from financial reports (text), quality control photos taken on a manufacturing line (images), or video captured inside the chemical reactors of an oil refinery.

In the second section, you will learn about data quality, data cleaning, and data provenance. It is important to understand potential problems with the data you use, how to identify them, and how to address them. Data provenance, that is, understanding where the data was collected or created, and how it was processed, is important because errors or biases may be introduced at various stages of the processing and handling pipeline.

The final section introduces different sources data sources. While most data for business analytics is internally produced by an organization, there is a vast amount of external information available to use and to combine with internal data for richer business analytics.

char	Individual Characters
string	A string of characters
byte	1 byte, -128 ... 127 or one ASCII characters
int (16 bit)	”Short”, Integer numbers, -32,768 ... 32,767
int (32 bit)	”Long”, Integer numbers, -2,147,483,648 ... 2,147,483,647
int (64 bit)	Integer numbers, -9,223,372,036,854,775,808 ... 9,223,372,036,854,775
float	Decimal numbers, 6 to 7 significant digits, ”single precision”
double	Decimal numbers, 15 to 16 significant digits, ”double precision”
boolean	Logical, true/false, 1 or 0

Table 2.1: Primitive Data Types

2.2 Data Types

2.2.1 Primitive Types

Primitive data types are basic types of data built into programming languages and other software systems such as statistics and analytics tools. They represent the simplest forms of data and serve as the building blocks for constructing more complex data structures.

Table 2.1 shows a list of common data types. However, not all software systems use the same names, and not all systems make the same distinctions. For example, the R system uses the terms `numeric` (which is actually a double type) and `integer` (which is a 32 bit integer).

Additionally, it is important in statistics and analytics to indicate the lack of a value, that is, a ”missing value”. Different systems use different special names for this. The R statistical system uses the term ”NA”, while the Python programming languages uses ”None” and the SQL database language uses the term ”Null”. Moreover, the meaning of these in practice can be ambiguous and says nothing about the reason for the missingness. For example, is the value not appropriate to the thing measured (e.g. in a table of geometric objects, the diameter value is simply not appropriate for two-dimensional object)? Was it missed during data collection? Was it withheld during data collection? Was it removed during initial analysis as an outlier?

Numbers

As Table 2.1 indicates, decimal numbers can be represented using different numbers of bytes with different precisions. Integer numbers are relatively straightforward. Here, a number is simply represented as its equivalent *binary number* (base 2, with digits 0 to 1), often with the first bit indicating the sign (positive/negative).

To represent decimal numbers, computers use the floating point representation defined

by the IEEE 754 standard¹. Figure 2.1 shows how decimal numbers are represented in binary form. A `float`, or single precision number occupies 4 bytes (32 bits): 1 bit for the sign, 8 bits for an exponent, and 23 bits for the fraction (also called "significand" or "mantissa"). With this, a `float` has a precision of approximately 6 to 7 decimal digits and a range at full precision between $\pm 1.18 \times 10^{-38} \dots \pm 3.4 \times 10^{38}$.

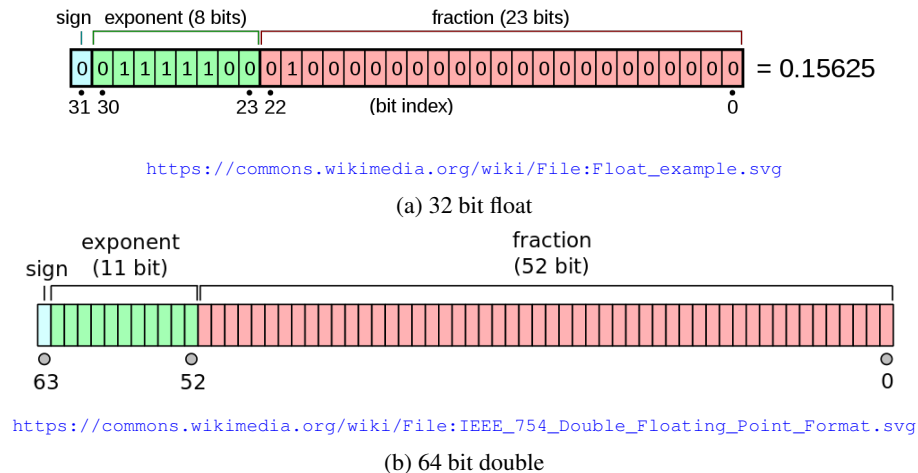


Figure 2.1: Floating Point Numbers (IEEE 754 Standard)

A `double`, i.e. a double precision number, occupies 8 bytes (64 bits): 1 bit for the sign, 11 bits for an exponent and 52 bits for the fraction. It has a precision of 15 to 16 decimal digits and a range at full precision between $\pm 2.23 \times 10^{-308}$... $\pm 1.80 \times 10^{308}$.

Example: Consider the number 0.15625 in the top of Figure 2.1. Its IEEE 754 float representation can be understood as follows:

1. First, convert 0.15625 to binary, which is 0.00101_2 ($1/8 + 1/32$).
 2. Rewrite in normalized scientific notation: $1.01_2 \times 2^{-3}$.
 3. Sign Bit: 0.15625 is positive, so the sign bit is 0.
 4. Exponent: The actual exponent is -3 , and the biased exponent is $-3 + 127 = 124$, which is 01111100_2 in binary.
 5. Fraction (significand, mantissa): The fraction is the normalized value without the leading 1, so it is $010000000000000000000000_2$ (23 bits).

Combining these components as shown in Fig. 2.1 leads to the number "01011111 00010000 00000000 00000000", written in 4 bytes of 8 bits each.

While the IEEE 754 standard defines how computers store decimal numbers internally, when exchanging information, numbers are printed as plain text. Such "printing as

¹https://en.wikipedia.org/wiki/IEEE_754

plain text” is called “*serialization*”, because the data are written as a series of characters or bytes. Writing out decimal numbers is fraught with complexities due to different idiosyncratic styles of writing or formatting numbers, depending on the application or the locale (that is, the dominant rules in the location of the user).

Format	Comment
-1023476.56	
-1023476,56	some locales use comma as decimal separator
-1,023,476.56	some locales use comma for grouping
-1.023.475,56	some locales use comma as sep and points to group
(1,023,476.56)	some applications use brackets for negation
- 1 023 476.56	some locales use space for grouping
-1.02347656e+06	“scientific notation”
-1023.47656e+03	also “scientific notation”

Table 2.2: Serializing Numbers to Text

The most frequent variations occur with respect to the decimal point (some locales, for example in Europe, use a comma instead), the grouping of digits (some locales group thousands, millions, etc. with spaces, points, commas, and other characters), writing negative numbers (in accounting, negatives are often put in parentheses instead of using a minus sign), and “scientific notation”, which specifies numbers as coefficients and exponent for powers of 10 (for example $1.234e+3 = 1.234 \times 10^3 = 1234$; $1.234e-2 = 1.234 \times 10^{-2} = 0.01234$). Table 2.2 shows examples of some idiosyncratic ways of writing the same number.

It is important to verify the number format in any data set, and to transform it into one that is readable and usable by the chosen business analytics tool.

Characters & Strings

There exist a multitude of writing systems beyond the Latin alphabet, using many different symbols. Symbols can represent consonants, consonant-vowel sequences, phonemes, words or morphemes, or syllables, leading to a vast range of symbols across the written languages of the world.



The Unicode system² was developed to address the limitations of earlier encoding systems and to enable consistent, universal representation of text from all the world’s writing systems. Before Unicode, there were different encoding systems, such as ASCII (American Standard Code for Information Interchange), which could only represent a limited set of characters (primarily used

²<https://home.unicode.org/>

in the English language). This led to difficulties in representing text in languages with larger character sets or different scripts. The Unicode Consortium was founded in 1988 and incorporated in 1991 with the goal of developing a universal character encoding standard. Unicode was standardized in 1998 as ISO/IEC standard 10646 and its popular UTF-8 encoding was standardized as RFC 2279³. Over the years, Unicode has been expanded and refined to include a wider array of characters, symbols, and scripts. This includes not only modern languages but also historic scripts, mathematical symbols, emojis, and more. By providing a unique identifier for every character, regardless of the computer system, software application, or programming language, Unicode solves the problem of inconsistent encoding and ensures that text appears consistently across different systems and devices. It has become the standard for modern software and internet protocols and is fundamental for web content, databases, applications, and more. The latest version of Unicode (v15.1) contains 149,813 characters for different 161 scripts, including 3782 emojis⁴.

UTF-8 (Unicode Transformation Format – 8-bit) is the most common method for representing Unicode characters. It uses between one and four bytes to represent a character and is backwards compatible with ASCII because the initial 127 Unicode characters are identical to the corresponding ASCII characters.

Example: Consider the word "Inuktitut". This word is written in the Inuktitut writing system as Δ¤ӮӰӮ, which contains six symbols (the Inuktitut writing system is not alphabetic, it is syllabic)⁵. The corresponding Unicode character numbers ("code-points") are: U+1403, U+14C4, U+1483, U+144E, U+1450, U+1466. These are given as *hexadecimal numbers*, using a base of 16 with digits from 0 to F. In text documents this may be written as \u1403 \u14c4 \u1483 \u144e \u1450 \u1466 when the document is read/parsed by an appropriate software tool that can understand this way of writing Unicode characters.

The corresponding decimal (base 10, with digits from 0 to 9) Unicode character numbers are 5123, 5316, 5251, 5198, 5200, 5222. These are used when the text is written for web content in HTML as HTML entities, such as "ᐃ ᓄ ᒃ ᑎ ᑐ ᑦ".

Using UTF-8, each of these six characters can be encoded in 3 bytes. These are usually written in hexadecimal form, indicated by the "0x" prefix. Hexadecimal form is a base 16 system and uses "numbers" between 0 and F, that is, it uses 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, A, B, C, D, E, and F as numbers. For example, 0xE1 is 14 (the "E") times 16 (the first digit) + 1 times 1 (the second digit), yielding 225. 0x.

The sequence of Unicode characters for the word "Inuktitut" becomes 0xE1 0x90 0x83 (first symbol) 0xE1 0x93 0x84 (second symbol) 0xE1 0x92 0x83 (third symbol) 0xE1 0x91 0x8E (fourth symbol) 0xE1 0x91 0x90 (fifth symbol) 0xE1 0x91 0xA6 (sixth symbol).

³<https://datatracker.ietf.org/doc/html/rfc2279>

⁴<https://www.unicode.org/versions/stats/>

⁵Using <https://www.inuktitutcomputing.ca/Transcoder/index.php> and <https://www.compart.com/en/unicode/>

As a business analyst, you may come across data that contains Unicode characters either spelled out in the "\uXXXX" form, or UTF-8 encoded in byte sequences, or in the HTML format. While you do not need to understand the technical details of Unicode and its different encodings, you should be aware that data in this format is common and you need to know how to deal with it when you encounter it. This includes using a Unicode-aware data storage and management system, using a Unicode-aware business analytics tool, and using a Unicode-aware visualization or report-writing tool.

Hands-On Exercise

- Choose your favourite emoji
- Determine its Unicode number ("codepage")
- Determine its UTF-8 encoding

Dates and Times

The world has largely standardized on the Gregorian calendar for secular and commercial use, while other calendar systems exist now only for religious or traditional purposes. However, as with written numbers, written dates show a bewildering variety of forms, depending on the locale and other traditions.

Complexities are introduced by 12 hour (AM/PM) versus 24 hour time formats ("14:30" is "2:30PM"), different time zones across the world, leap seconds and leap years, week numbering (does it start with the first full week?), different written formats for the sequence of days, months, and years (is "06-07-09" June 7, 2009 or July 6, 2009, or July 9, 2006?), different separators between years, dates, and months ("06/07/09" and "06-07-09"), and the difficult arithmetic when using years, months, and days.

The ISO 8601⁶ (first published in 1988) and RFC 3339⁷ (published in 2002) standards define how dates and times should be written. Table 2.3 shows a summary of the ISO 8601/RFC3339 rules (the italicized forms are in ISO 8601 but not in RFC 3339). However, these standards are by no means universally accepted and reading/parsing date and time data remains a difficult and complex task in many business analytics settings.

Even within the ISO 8601 standard, there are numerous ways to express the same data or time, such that June 13, 2024 can be written as an ordinal date ("2024-165") or a week date ("2024-W24-4") and the time of T13:45:30 can be written as "T13:45.500" or as "T1345.500".

For a business analyst, it is important to verify the format of dates and times in any data set, especially when the data comes from different or external sources.

⁶https://en.wikipedia.org/wiki/ISO_8601

⁷<https://datatracker.ietf.org/doc/html/rfc3339>

Calendar dates	YYYY-MM-DD
<i>Ordinal dates</i>	YYYY-DDD
<i>Week dates</i>	YYYY-Www-d
Times	<i>Thh:mm:ss.sss (or Thhmss.ss)</i> <i>Thh:mm:ss (or Thhmss)</i> <i>Thh:mm.mmm or Thmm.mmm</i> <i>Thh:mm or Thmm</i> <i>Thh.hhh</i>
Time Zones	<i><time>Z or <time>±hh:mm or</i> <i>(<time>±hhmm or <time>±hh)</i>
Combined	<i><date>T<time></i>
Periods	<i>PnYnMnDTnHnMnS or P<date>T<time></i>

The *italicized* forms are in ISO 8601 but not in RFC 3339

Table 2.3: ISO 8601 / RFC 3339 Rules for Dates and Times

A year is a leap year if it can be divided by four and (cannot be divided by 100 but can be divided by 400). Formally:

```
(year % 4 == 0) and (year % 100 != 0 or year % 400 == 0)
```

Hands-On Exercise

The territory of Nunavut was created on April 1st, 1999.

- Express the date in RFC 3339
- Calculate the number of days since the creation of Nunavut
- Assume that a ceremony took place at 3PM that day in Iqaluit and express this date-time in RFC 3339
- Assume the ceremony lasted for 125 minutes and express this duration in RFC 3339

Collections

Collections can store multiple instances of primitive data, often heterogeneous. Different collection types have different characteristics in terms of whether they are

- ordered or unordered,
- homogeneous (same primitive types) or heterogeneous (different primitive types),
- unique or allow duplicates,

Python		
list	[1, 2, "a", "b", 2]	mutable, ordered
tuple	(1, 2, "a", "b", 2)	immutable
set	{1, 2, "a", "b"}	mutable, unordered, unique
dict	{"make": "Ford", "year": 2023}	mutable

R		
list	list(1, 2, "a", "b", 2)	mutable, ordered
vector	c(1, 2, 3)	mutable, same primitive type
factor	as.factor(c("Hot", "Med", "Cold"))	ordered
matrix	matrix(c(1, 2, 3, 4), nrow=2)	
array	array(c(1, 2, 3), c(4, 5, 6))	

Table 2.4: Structured data types ("collection types") in Python and R

- mutable (can be changed) or immutable (cannot be changed).

Different software tools offer different kinds of structured types and unfortunately the terminology is not necessarily consistent across software tools. Table 2.4 provides a summary of structured types in Python and R, with examples and key characteristics of the type.

2.2.2 Structured Data

Data that you encounter in business analytics is built on the primitive and collection data types described in the previous section. We distinguish between structured data, such as tables, key-value pairs, documents, or graphs, and unstructured data, such as text, images, and audio/video data. The latter are called unstructured because these data essentially come as simply a sequence of characters or bytes. Information must first be identified in them and extracted from them, before it can be used for analytics.

Tables

Table data refers to a method of organizing data in a structured, tabular format, where the data is arranged in rows and columns. Each *row* in a table represents a single record or entry. For instance, in a table of customer data, each row could represent a different customer. *Columns*, sometimes called *fields*, represent different attributes or characteristics or features of a record. In the customer data example, columns might represent attributes of a customer such as their name, address, and purchase history. The intersection of a row and a column is called a cell. Each *cell* contains a single piece of data for a particular attribute of a record. For example, the cell at the intersection of the "Name" column and the third row might contain the name of the third customer. Cells may be of simple type or be themselves of structured types, such as sets or lists or even other tables. Tables often have a *header row* at the top, which contains the names of the columns. These headers provide context for what each column in the table represents. Table 2.5 shows an example table with a header row that names the columns, three rows of data in three columns that use simple data types (strings and

integers). Some tables may also have an *index column* as the first column with row numbers.

CSV Files While table data is familiar from spreadsheet systems such as Microsoft Excel or LibreOffice Calc, these tools often store table data in a format that is unique to their system and difficult to read with other tools. This is because spreadsheet tables can contain formulas, formatting instructions such color and font, and other information besides the actual data.

The standard format for storing and exchanging tabular data (i.e. the *serialization format*) is the comma-separated value file ("CSV" file) that is standardized in RFC 4180⁸. Tabular data is stored in a plain text format, without formatting instructions or formatting information, that makes it easy to read and write with different software tools, such as statistics and analytics software, spreadsheet applications, or database management systems.

CSV files are plain text files, typically encoded in ASCII or UTF-8. Every line contains one row of the table, and fields within a row are separated by commas (although sometimes other, non-standard delimiters such as semicolon are used). Fields are typically of a primitive data type although the interpretation of the field content is left to the software tool reading the CSV file. The CSV file may contain an optional header as the first line, with the same format as the data lines of the file. Every line is ended by a line break using the sequence of CR and LF characters⁹. Every line must contain the same number of fields and fields are allowed to be empty, but must still be separated by a comma. The content of each field may be enclosed by double quotes (although sometimes other, non-standard quotes like single quotes are used). The following is a CSV serialization of Table 2.5:

```
"Name", "Area", "Population" CR LF
"Canada", "9984670", "38781292" CR LF
"Nigeria", "923768", "223804632" CR LF
"Germany", "357600", "83294633" CR LF
```

⁸<https://datatracker.ietf.org/doc/html/rfc4180>

⁹These stand for "carriage return" and "line feed", respectively, and are a hold-over from the era of mechanical typewriters where the paper carriage needed to be returned to the start of a line and then advanced by one line. CR and LF are represented by ASCII/UTF-8 codes 13 and 10, respectively. Hence, the CSV line break conforms to the Microsoft Windows convention of line breaks. MacOS and Linux use only the LF character for line breaks.

Name	Area	Population
Canada	9,984,670	38,781,292
Nigeria	923,768	223,804,632
Germany	357,600	83,294,633

Table 2.5: Example Table

While the CSV format is standardized, not all data sets necessarily conform fully to the standard. You may encounter different field delimiters, such as semicolons, tabs, carets ("^") or others. Line breaks may not use the Microsoft Windows convention of CR LF but instead use only the LF character as is typical on MacOS and Linux/Unix systems. Not all fields may be quoted and you may encounter a mix of double quotes and single quotes even in the same CSV file. Additionally, the field contents themselves, such as numbers and dates, may themselves not be standards compliant and exhibit a range of different notations, as discussed above. For a business analyst, it is important to recognize these variations and be able to address them prior to further data analysis.

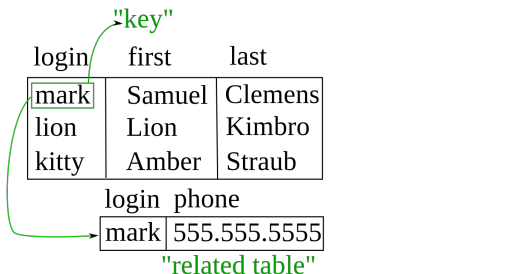
Hands-On Exercise

- Search the internet for a CSV file of the population and areas of all countries of the world
- Examine the CSV file and answer the following questions:
 - What is the delimiter?
 - Which fields are quoted, and how?
 - What is the line ending character(s)?
 - What is the number format?
 - What is the date format (if there are dates)?
- Import the CSV file into your favourite spreadsheet tool
 - Does it recognize all information correctly? If not, what is not imported well?
- Export the CSV file from your tool under a different name.
 - Do you get an identical file to the one you imported? If not, what has changed?

Relational Databases Tabular data is also the basis for relational database management systems (RDBMS). Tables in these systems are called relations¹⁰. Records, i.e. rows of a relation, are uniquely identified by *primary keys*. These may be "natural" primary keys, such as a combination of fields (also called "attributes" in RDBMS) or artificial/synthetic primary keys. For example, in some applications one may assume that the combination of first name, last name, date of birth, and postal code uniquely identifies a person and is used as a primary key. However, it is generally safer to assign artificial primary keys, such as consecutive numbers, to records.

One key characteristic of data in an RDBMS is that fields in one table can refer to primary key fields in another table. For example, the product numbers for an order in the order table must refer to product numbers of products in the products table. The referring fields are called "*foreign keys*". Foreign-key relationships ensure *referential integrity*, a form of validity of the data. They also allow an RDBMS to easily retrieve related records from different relations. Figure 2.2 shows an example of keys in a relational database.

¹⁰After the mathematical concept of a relation as a subset of a cross-product.



https://commons.wikimedia.org/wiki/File:Relational_key_SVG.svg

Figure 2.2: Keys in a relational database

Data *normalization* in an RDBMS refers to reducing data redundancy. For example, if a customer can have multiple addresses, rather than using multiple address fields in the customer relation or having multiple customer records for a customer (one for each address), normalization will create a table to store addresses where each address refers back to a particular customer using a foreign-key relationship. Normalizing relations and thereby reducing redundancy makes data storage more efficient and reduces the potential for inconsistent data, leading to higher data integrity. RDBMS typically use the structured query language (SQL) for retrieving information.

Prominent RDBMS examples are Oracle RDBMS¹¹, a proprietary system for on-premises installation; the PostgreSQL¹² open-source RDBMS system for on-premises installation; and Amazon RDS¹³, Google BigQuery¹⁴, and Azure SQL¹⁵ which are cloud-based systems on AWS, Google Cloud, and Microsoft Azure, respectively.

Hands-On Exercise

- Assume that products are identified by a product code and have attributes such as description, weight, and price.
- Assume that suppliers are identified by a supplier number and have attributes such as name and address.
- Assume that each product is available from exactly one supplier (but a supplier can supply multiple products).

Write example relations and identify foreign-key relationships for referential integrity, similar to Figure 2.2

¹¹<https://www.oracle.com/ca-en/database/>

¹²<https://www.postgresql.org/>

¹³<https://aws.amazon.com/rds/>

¹⁴<https://cloud.google.com/bigquery>

¹⁵<https://azure.microsoft.com/en-ca/products/azure-sql/database>

Key-Value Data Stores

Key-value data stores are a type of non-relational (NoSQL¹⁶) database that organize data as a collection of key-value pairs. In this model, each data item is stored as a key, along with an associated value. Keys may have multiple components, in an ordered list of "minor keys". The associated value is not interpreted by the data store, and can contain anything that is meaningful to the application, from primitive data types to collections to complex documents to images or video data. Figure 2.3 shows an example of the key-value model of data storage.

Key	Value
K1	AAA,BBB,CCC
K2	AAA,BBB
K3	AAA,DDD
K4	AAA,2,01/01/2015
K5	3,ZZZ,5623

<https://commons.wikimedia.org/wiki/File:KeyValue.PNG>

Figure 2.3: Key Value Data Store

Important characteristics of key-value stores the extremely simple data model: every item is stored as a key and its corresponding value. The keys are unique identifiers. Due to their simple structure, key-value databases allow faster data insertion, updating, and retrieval when compared to more complex relational databases. Unlike relational databases, key-value stores do not have predefined relations with foreign-key relationships. This means that the values associated with keys can be changed dynamically, and different keys can have values of different types. This makes key-value stores more flexible than other databases. On the other hand, they lose the data integrity advantages that come from a predefined schema and the referential integrity based on relationships between multiple tables. Key-value stores are more efficient at storing information than RDBMS because empty table cells do not need to be stored. Key-value stores are also easier to scale and distribute among multiple computers, due to their simple data model. On the other hand, key-value stores are limited in terms of their data querying and analysis capabilities. They are not inherently designed for complex queries, such as joining data across different keys.

Example key-value data stores include Redis¹⁷, an open-source, in-memory key-value store. Amazon DynamoDB¹⁸ is a proprietary scalable NoSQL database service available on the AWS cloud. Google BigTable¹⁹ and Azure CosmosDB²⁰ are key-value stores offered on the Google cloud and the Microsoft Azure cloud. Facebook's RocksDB,

¹⁶NoSQL is term to describe non-relational database models. It does not mean "no SQL", but means "Not only SQL."

¹⁷<https://redis.io/>

¹⁸<https://aws.amazon.com/dynamodb/>

¹⁹<https://cloud.google.com/bigtable>

²⁰<https://cosmos.azure.com/>

Google's LevelDB²¹ and the Apache Cassandra²² and HBase²³ projects offer open-source systems for on-premises installation.

Documents (JSON)

When speaking about documents in the context of structured data, we do not mean unstructured text (as a series of characters) but a structured collection of elements. The JavaScript Object Notation (JSON) is a lightweight data-interchange format that is easy for humans to read and write, and also easy for machines to parse and generate. It is software tool independent. Originally developed for exchanging data between web servers and web browser client applications, it has emerged as a popular way of describing and exchanging many different kinds of data for a variety of purposes in many different applications. JSON was standardized in RFC 8259²⁴ in 2017.

JSON documents are plain text documents encoded in UTF-8 and consist of key-value pairs where the key or name is a string and the separator is a colon. Values may be strings (enclosed by single or double quotes), numbers, boolean values ("true" or "false"), or the special value "null". Values are either objects or arrays. *JSON objects* are unordered collections of zero or more key-value pairs and are delimited by "{" and "}". Figure 2.4 shows an example of a JSON object with key-value pairs and nested objects. *JSON arrays* are ordered sequences of zero or more values and are delimited by "[" and "]". Arrays contain values but no keys and, because they are ordered, elements can be accessed by position. Figure 2.5 shows an example of a JSON array, i.e. a list of values, in this example a list of objects.

```
{
  "Image": {
    "Width": 1060,
    "Height": 400,
    "Title": "Skyline of Iqaluit, Nunavut",
    "Url": "https://upload.wikimedia.org/wikipedia/commons/b/b4/Iqaluit_skyline.jpg",
    "Legal": {
      "Copyrighted": true,
      "License": "GNU Free Documentation License",
      "Inception": "2010-03-24",
      "Author": "Aaron Lloyd"
    }
  }
}
```

Figure 2.4: JSON Example – Complex Object

²¹<https://github.com/google/leveldb>

²²<https://cassandra.apache.org/>

²³<https://hbase.apache.org/>

²⁴<https://datatracker.ietf.org/doc/html/rfc8259>

```
[  
  {  
    "Latitude": 56.536389,  
    "Longitude": -61.718889,  
    "City": "Nain",  
    "Province": "NL",  
    "Postal": "A0P",  
    "Country": "Canada"  
  },  
  {  
    "Latitude": 53.512778,  
    "Longitude": -60.135556,  
    "City": "Sheshatshiu",  
    "Province": "NL",  
    "Postal": "A0P",  
    "Country": "Canada"  
  }  
]
```

Figure 2.5: JSON Example – List of Objects

Hands-On Exercise

Describe yourself in a JSON object:

- Identify information about yourself, such as names, addresses, dates, relationships (work, school, uni), etc.
- Structure the information in JSON Objects and Arrays
- Use nested structures, e.g. objects in arrays, or arrays in objects, or objects in objects, etc.

Documents (XML)

XML²⁵, for “eXtensible Markup Language”, is a flexible and versatile serialization format that plays an important role in the storage and transmission of data. It is a text-based format that allows for the creation of custom tags (tags are used to describe and delimit data elements), providing a means to define and structure data in a way that is both machine-readable and human-readable. Unlike HTML (the Hypertext Markup Language that describes web pages), which has a predefined set of tags for web page layout, XML does not prescribe any specific tags, allowing users to create tags tailored to their specific application.

XML’s development began in the late 1990s by an XML Working Group under the auspices of the World Wide Web Consortium (W3C). XML 1.0 was officially recommended by the W3C in February 1998. It quickly gained widespread adoption due to its simplicity, extensibility, and ability to work seamlessly across different systems and

²⁵ <https://www.w3.org/TR/2008/REC-xml-20081126/>

platforms. XML has become a cornerstone technology in numerous domains, from web services and APIs to configuration files and data interchange formats.

An XML document is composed of *elements*. XML elements are described by matching opening and closing *tags*, between which simple text content or other XML elements may be placed. Elements in turn may have *attributes*. Attributes are specified in the opening tag of an element and may contain simple, quoted text data only. This hierarchical organization can represent complex data relationships and nested structures.

XML files are human-readable and self-descriptive in nature; the names of elements and attributes used in the document indicate that type or meaning of the data they describe, enhancing the understanding and interpretation of the data's structure and meaning. Additionally, XML is platform-independent and language-neutral, making it a universally accepted standard for data interchange across different systems and applications.

XML element and attribute names may be defined within a *namespace*. This allows mixing elements with the same name but different namespaces in the same document and removes ambiguity that could arise from identically named elements that describe different data or content. For example when mixing customer and product information in an order document, both the customer and the product may have a "name" element and different namespaces for the two "name" elements helps to tell them apart. Namespaces are declared using the special `xmlns` attribute and are typically defined by a URI (Uniform Resource Identifier), typically a URL (Uniform Resource Locator). These URI/URL are for identification purposes and do not need to describe an actually existing resource.

The following example describes the Innu people of northern Canada in the form of an XML document. The element names, like `People`, `History`, `Culture` are self-descriptive and human-readable. Note that each opening tag (such as `<Traditions>`) is matched by a corresponding closing tag (such as `</Traditions>`). Empty elements that do not contain any content are defined using a single tag (for example, the `<geo:Location.../>` element). Some elements contain attributes, such as the `Name` attribute of the `GeneralInformation` element. Attributes must be quoted character strings.

Namespaces are declared at the root element of the document: The `xmlns:geo` and `xmlns:hist` are namespace declarations. They are used to distinguish between geographical (`geo`) and historical (`hist`) data. Notice how element and attribute names may be prefixed by a namespace (such as the `<hist:History>` element or the `geo:Country` attribute. The `xmlns` declaration defines the default namespace that applies to all elements and attributes without an explicit namespace.

Notice that elements with the same name may be repeated. For example, there are multiple `Period` elements in the `History` element, each with their own `hist:era` attribute to specify the era. This allows one to represent lists or sets of elements.

```

<People
    xmlns="https://www.example.com/peoples"
    xmlns:geo="http://www.example.com/geo"
    xmlns:hist="http://www.example.com/history">
    <GeneralInformation>
        Name="Innu" Language="Innu-aimun">
            <geo:Location geo:Country="Canada"
                geo:Regions="Labrador, Quebec" />
        </GeneralInformation>
        <hist:History>
            <hist:Period hist:era="Pre-Colonial">
                <Description>
                    Nomadic lifestyle, primarily
                    hunting and fishing.
                </Description>
            </hist:Period>
            <hist:Period hist:era="Post-Colonial">
                <Description>
                    Impact of colonization,
                    including displacement and
                    cultural changes.
                </Description>
            </hist:Period>
        </hist:History>
        <Culture>
            <Traditions>
                <Tradition>
                    Hunting and fishing as cultural
                    and subsistence activities.
                </Tradition>
                <Tradition>
                    Use of the tepee for temporary
                    shelter.
                </Tradition>
            </Traditions>
            <Art>
                <Form>Drum making</Form>
                <Form>Clothing with intricate beadwork
                </Form>
            </Art>
        </Culture>
        <Challenges>
            Issues like land rights, cultural preservation
        </Challenges>
    </People>

```

Given the similarities between JSON and XML it is not surprising that one can readily be transformed into the other. An equivalent JSON document of the above XML document could be as follows:

```

" Innu": {
    "@xmlns:geo": "http://www.example.com/geo",
    "@xmlns:hist": "http://www.example.com/history",
    "GeneralInformation": {
        "@Name": "Innu",
        "@Language": "Innu-aimun",
        "Location": {
            "@geo:Country": "Canada",
            "@geo:Regions": "Labrador, Quebec"
        }
    },
    "History": [
        "Period": [
            {
                "@hist:era": "Pre-Colonial",
                "Description": "Nomadic lifestyle,  
primarily hunting and fishing."
            },
            {
                "@hist:era": "Post-Colonial",
                "Description": "Impact of colonization,  
including displacement and  
cultural changes."
            }
        ]
    ],
    "Culture": {
        "Traditions": [
            "Tradition": [
                "Hunting and fishing as cultural and  
subsistence activities.",
                "Use of the tepee for temporary shelter."
            ]
        ],
        "Art": [
            "Form": [
                "Drum making",
                "Clothing with intricate beadwork"
            ]
        ]
    },
    "CurrentStatus": {
        "Challenges": "Issues like land rights,  
cultural preservation, etc."
    }
}

```

In this example, XML namespaces are represented as properties with names prefixed by "@". This does not imply any special meaning or treatment in JSON, but makes it easier for the computer to read and parse ("understand") the document. XML elements with attributes are represented as JSON objects, while repeated elements are represented as JSON arrays.

In comparing XML to JSON, it is evident that both formats are human as well as

machine readable. It is also clear that XML is more verbose or lengthy. This is an advantage in that it makes it very self-descriptive, but a disadvantage in that XML documents are larger than corresponding JSON documents. In contrast, JSON is more compact or lightweight, and not quite as self-descriptive as an XML document. XML supports more complex structures than JSON through its attributes, namespaces, and a larger selection of possible data types for simple content.

While XML can be strictly defined using XML Schema there is not yet a well-adopted means for specifying JSON documents. This means that JSON documents cannot be validated against a set of rules or constraints, possibly leading to data quality issues, but, on the other hand, may be used more flexibly.

Hands-On Exercise

Describe yourself in an XML document:

- Identify information about yourself, such as names, addresses, dates, relationships (work, school, uni), etc.
- Structure the information in Elements and Attributes
- Use nested elements where appropriate

Document Databases Document databases, a type of NoSQL databases, are designed to store, retrieve, and manage document-oriented information, typically in the form of JSON or BSON (Binary JSON) documents. Unlike traditional relational databases that store data in rows and columns, document databases handle data in a more flexible, semi-structured way. They are designed for handling large volumes of diverse data that does not fit into a tabular format. Document databases may be thought of as nested key-value data stores where all keys are strings. Their fundamental unit of storage is the document. Unlike relational databases, document databases usually do not have a predefined schema. Each document in a collection can have its own unique structure, with different fields, data types, and sizes. This makes them more flexible than RDBMS. Document databases often offer query languages that are designed to handle complex queries on document data, including searching within documents and aggregating data across multiple documents.

Typical use cases for document databases are content management, where different types of content need to be stored and retrieved efficiently, catalogs and product data, where each product may have different attributes and structures, and real-time analytics of Internet-of-Things (IoT) sensor data, where large volumes of unstructured and semi-structured data are generated.

Prominent examples of document databases are MongoDB²⁶ and ArangoDB²⁷ which are partially proprietary system for on-premises installation or cloud-based use; Apache

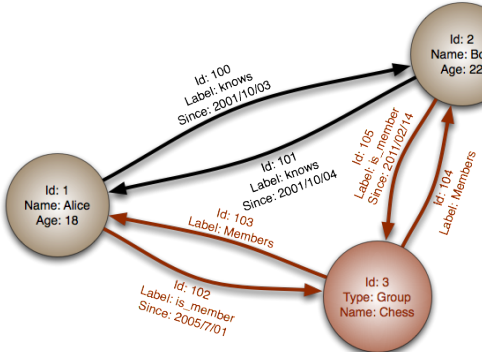
²⁶<https://www.mongodb.com/>

²⁷<https://arangodb.com/>

CouchDB²⁸ which is a fully open-source system; and AWS DocumentDB²⁹ which is a cloud-based system on the Amazon AWS cloud.

Graphs

Graphs consist of *nodes* (also called *vertices*) and *edges* (also called *arcs* or *relationships*) that connect two nodes. Edges may be directed or undirected. Both nodes and edges may be labelled or typed. For example, different node types may be used to represent customers and products; different edge types may represent a customer ordering a product, a customer returning a product, a customer obtaining a quote about a product, etc. Graph data is found in social networks (between people, events, topics, etc.), in logistics networks (between suppliers, customers, warehouses, distribution centers, etc.), in financial networks (between organizations, accounts, etc.), in biological networks, and in many other contexts.



https://commons.wikimedia.org/wiki/File:GraphDatabase_PropertyGraph.png

Figure 2.6: Property Graph Example

Graph data is either in the form of property graphs or RDF graphs ("Resource Description Framework"). *Property graphs* are a graph data model where each node and edge can have a set of properties (key-value pairs) that describe the attributes of the entity represented by the node. Edges can also have properties, which can describe attributes of the relationship. For example, a node representing a person might have properties like `name: "John Doe"` and `age: 30`. An edge representing a friendship relationship might have a property like `since: 2010`. Property values may be simple data types or complex ones like JSON documents. Figure 2.6 shows an example property graph.

In contrast to property graphs, RDF³⁰ graphs do not allow properties on nodes and edges. Instead, they describe information in subject–predicate–object triples. What might be an "age" property in a property graph can be described in RDF as the triple

²⁸<https://couchdb.apache.org/>

²⁹<https://aws.amazon.com/documentdb/>

³⁰<https://www.w3.org/RDF/>

JohnDoe – has age – 30. In an RDF graph, subjects and objects have unique identifiers (URIs, uniform resource identifiers³¹) that are typically defined in the form of URL as defined in RFC 2616³²), or are literal values such as strings or numbers. Figure 2.7 shows an example RDF graph.



Figure 2.7: RDF Graph Example

While graphs could be modelled or described in table (relational) form, or as key-value pairs, graph databases provide powerful, intuitive, and efficient graph-specific queries³³. Figure 2.8 shows an overview of different query types:

- *Path queries*: Reachability of nodes, shortest-path between nodes
- *Subgraph queries*: Exact or approximate match of a smaller graph in a larger one
- *Aggregate queries*: Aggregating nodes or properties along paths
- *Similarity search*: Similarity of nodes or edges using path-based approaches, graph embedding-based approaches
- *Keyword search*: Tree-based semantics, subgraph-based semantics
- *Natural language query answering*: Identifying edges or nodes

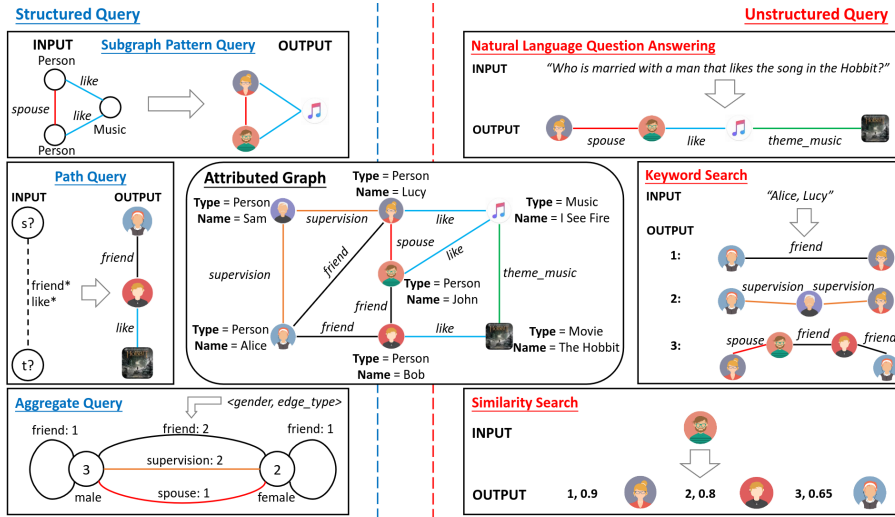
Prominent examples of graph database management systems are JanusGraph³⁴, a fully

³¹<https://www.w3.org/TR/webarch/#identification>

³²<https://www.ietf.org/rfc/rfc2616.txt>

³³Source: Wang, Y., Li, Y., Fan, J., Ye, C., & Chai, M. (2021). A survey of typical attributed graph queries. World Wide Web, 24, 297-346

³⁴<https://janusgraph.org/>



Source: Wang, Y., Li, Y., Fan, J., Ye, C., & Chai, M. (2021). A survey of typical attributed graph queries. World Wide Web, 24, 297-346.

Figure 2.8: Graph Queries

open-source system for on-premises installation; Arangodb³⁵, Neo4J³⁶, and OrientDB³⁷ which are partially open-source for on-premises installation; AWS Neptune³⁸ on the Amazon AWS cloud, and CosmosDB³⁹ on Microsoft Azure cloud.

While there does not exist a standardized serialization format for property graph data interchange, PG-JSON and GraphSON are two recent proposals to describe property graphs in a JSON object, shown in Figure 2.9 and Figure 2.10.

³⁵<https://arangodb.com/>

³⁶<https://neo4j.com/>

³⁷<https://orientdb.org/>

³⁸<https://aws.amazon.com/neptune/>

³⁹<https://azure.microsoft.com/en-ca/products/cosmos-db>

```
{  
  "nodes": [  
    {  
      "id": 101,  
      "labels": ["Person"],  
      "properties": {"name": ["Alice"], "age": [15], "country": ["USA"]}  
    },  
    {  
      "id": 102,  
      "labels": ["Person", "Student"],  
      "properties": {"name": ["Bob"], "country": ["Japan", "Germany"]}  
    }  
  ],  
  "edges": [  
    {  
      "from": 101,  
      "to": 102,  
      "undirected": true,  
      "labels": ["sameSchool", "sameClass"],  
      "properties": {"since": [2012]}  
    },  
    {  
      "from": 102,  
      "to": 101,  
      "labels": ["likes"],  
      "properties": {"since": [2015]}  
    }  
  ]  
}
```

Figure 2.9: PG-JSON Example

```
{  
  "graph": {  
    "mode": "NORMAL",  
    "vertices": [  
      {  
        "name": "lop",  
        "lang": "java",  
        "_id": "3",  
        "_type": "vertex"  
      },  
      {  
        "name": "vadas",  
        "age": 27,  
        "_id": "2",  
        "_type": "vertex"  
      },  
      {  
        "name": "marko",  
        "age": 29,  
        "_id": "1",  
        "_type": "vertex"  
      },  
      {  
        "name": "peter",  
        "age": 35,  
        "_id": "6",  
        "_type": "vertex"  
      },  
      ...  
    ],  
    "edges": [  
      {  
        "weight": 1,  
        "_id": "10",  
        "_type": "edge",  
        "_outV": "4",  
        "_inV": "5",  
        "_label": "created"  
      },  
      {  
        "weight": 0.5,  
        "_id": "7",  
        "_type": "edge",  
        "_outV": "1",  
        "_inV": "2",  
        "_label": "knows"  
      },  
      {  
        "weight": 0.400,  
        "_id": "9",  
        "_type": "edge",  
        "_outV": "1",  
        "_inV": "3",  
        "_label": "created"  
      },  
      ...  
    ]  
  }  
}
```

Figure 2.10: GraphSON Example

In contrast, multiple standardized RDF graph serializations are defined⁴⁰. The "Turtles" format (Terse RDF Triples) is the most compact representation and most easy to read for humans. The following example describes the RDF graph in Figure 2.7. It defines three prefixes of URIs to identify resources. These resources form the subjects, predicates, and objects of the RDF triples.

```
@prefix eric: <http://www.w3.org/People/EM/contact#> .
@prefix contact: <http://www.w3.org/2000/10/swap/pim/contact#> .
@prefix rdf: <http://www.w3.org/1999/02/22-rdf-syntax-ns#> .

eric:me contact:fullName "Eric Miller" .
eric:me contact:mailbox <mailto:e.miller123(at)example> .
eric:me contact:personalTitle "Dr." .
eric:me rdf:type contact:Person .
```

The equivalent N-Triples representation is still reasonably compact, but easier for computers to read and parse. Here, the URI prefixes are embedded in the subjects, predicates, and objects:

```
<http://www.w3.org/People/EM/contact#me>
<http://www.w3.org/2000/10/swap/pim/contact#fullName>
"Eric Miller" .

<http://www.w3.org/People/EM/contact#me>
<http://www.w3.org/2000/10/swap/pim/contact#mailbox>
<mailto:e.miller123(at)example> .

<http://www.w3.org/People/EM/contact#me>
<http://www.w3.org/2000/10/swap/pim/contact#personalTitle>
"Dr." .

<http://www.w3.org/People/EM/contact#me>
<http://www.w3.org/1999/02/22-rdf-syntax-ns#type>
<http://www.w3.org/2000/10/swap/pim/contact#Person> .
```

Finally, the RDF/XML serialization uses the XML language to describe RDF triples. It is quite verbose:

⁴⁰Examples taken from https://en.wikipedia.org/wiki/Resource_Description_Framework

```

<?xml version="1.0" encoding="utf-8"?>
<rdf:RDF xmlns:contact="http://www.w3.org/2000/10/swap/pim/contact#"
    xmlns:eric="http://www.w3.org/People/EM/contact#"
    xmlns:rdf="http://www.w3.org/1999/02/22-rdf-syntax-ns#">
    <rdf:Description rdf:about="http://www.w3.org/People/EM/contact#me">
        <contact:fullName>Eric Miller</contact:fullName>
    </rdf:Description>
    <rdf:Description rdf:about="http://www.w3.org/People/EM/contact#me">
        <contact:mailbox rdf:resource="mailto:e.miller123(at)example"/>
    </rdf:Description>
    <rdf:Description rdf:about="http://www.w3.org/People/EM/contact#me">
        <contact:personalTitle>Dr.</contact:personalTitle>
    </rdf:Description>
    <rdf:Description rdf:about="http://www.w3.org/People/EM/contact#me">
        <rdf:type rdf:resource="http://www.w3.org/2000/10/swap/pim/contact#Person"/>
    </rdf:Description>
</rdf:RDF>

```

Hands-On Exercise

Document yourself in a Turtle:

- Identify information about yourself, such as names, addresses, dates, relationships (work, school, uni), etc.
- Structure the information in Turtle triples
- Make up appropriate prefixes and appropriate verbs/predicates

2.2.3 Unstructured Data

Text

Text refers to written language in some writing system and is provided as a string of characters or a file containing bytes that encode the text in Unicode UTF-8 or some other format. Text data in business analytics does not normally contain formatting instructions, such as font sizes or font styles, or mixed tables or images, as might be common in word processing systems. Instead, it refers to plain text only.

Text analysis is the process of extracting meaningful information from unstructured text data. Typical text analysis tasks are named entity recognition, co-reference analysis, event extraction, sentiment analysis, and document clustering. Named entity recognition identifies names of persons, organizations, or places, and expressions of time, quantity, or monetary amounts in a text. It is useful for content classification and data extraction.

Co-reference analysis involves identifying when two or more expressions in a text refer to the same entity. This task is crucial for understanding the context and for maintaining the continuity of subjects throughout the text. For example, in the sentence "Alice drove her car. She parked it near the mall," co-reference analysis links "She" to "Alice"

and "it" to "Alice's car." Co-reference analysis helps in understanding the text flow and the relationships between various entities.

Event and relationship extraction is about identifying instances of specific types of events in text and the entities associated with them. An event can be anything that happens or is described as happening. For example, in "The company acquired a startup for \$1 million in 2021," event extraction would identify the acquisition event, involving the company, the startup, the amount of \$1 million, and the time 2021. This task is useful for information monitoring or historical data analysis.

Sentiment analysis involves identifying and categorizing opinions expressed in a piece of text, especially to determine whether the writer's attitude towards a particular topic, product, etc., is positive, negative, or neutral. It is widely used in social media monitoring, brand monitoring, customer service, and market research.

Document clustering is a method to categorize documents into groups (or clusters) based on their similarity. It is useful for news aggregation, organizing web search results, discovering prevalent topics or themes and grouping of similar documents to make it easier to find relevant information.

The history of text mining approaches has evolved through several stages. In the 1950s and 1960s, text mining began with symbolic approaches, involving rule-based systems. These systems, which relied on handcrafted linguistic rules, attempted to encode human language knowledge into a format readable by computers. Their reliance on extensive domain knowledge and manual rule creation made them inflexible and unable to adapt to language variations and new data.

The late 1980s and 1990s saw a shift towards statistical methods in text mining, driven by the growing availability of digital text data and computational power. This period was characterized by the use of machine learning models like Naive Bayes, Decision Trees, and Support Vector Machines. The era of corpus linguistics also emerged, enabled by the availability of large text corpora, allowing for the statistical analysis of real-world text data. Techniques like Latent Semantic Analysis (LSA) and Latent Dirichlet Allocation (LDA) were developed for topic modeling and document classification.

The 2010s marked a revolution in text mining with the advent of deep learning and neural networks, which provided the ability to learn complex patterns in large datasets. Recurrent Neural Networks (RNNs) and variants like LSTMs became popular for handling sequential text data. The development of attention mechanisms and transformer models in the 2020s, such as Google's BERT or OpenAI's ChatGPT, represented yet another significant advancement.

Hands-On Exercises

1. Identify a specific business problem that can be addressed by analyzing text data
2. What text data would you need to address the problem?
3. What would you wish to do with the text data?
4. Where might you get this text data?

Regular Expressions (RegEx)

Regular expressions (often abbreviated as regex or regexp) are a tool for pattern matching within text. They enable the specification of complex search patterns in a concise and flexible manner. Regular expressions are widely used for searching, editing, or manipulating text and data. For example, the following regular expression matches any number, including one in scientific notation⁴¹:

```
[+-] ? (\d+ (\.\d*)? | \.\d+) ([eE][+-]? \d+)? .
```

Metacharacter	Description
^	Matches start of text
.	Matches any character; matches the dot character when used within brackets
[]	Matches any of the characters in the brackets; - can be used to specify ranges of characters
[^]	Matches any character not in the brackets
\$	Matches the end of text
()	Marked subexpression
\n	Matches the n-th marked subexpression
*	Matches the preceding element zero or more times
{m,n}	Matches the preceding element at least m and not more than n times

Adapted from https://en.wikipedia.org/wiki/Regular_expression

Table 2.6: Basic Regular Expressions

Regular expressions are specified using *metacharacters*, i.e. characters that describe other characters. Table 2.6 shows the metacharacters for basic regular expressions as defined by the POSIX standard. All other characters are treated as literal characters. With these definitions, you can understand the examples shown in Table 2.7.

Extended regular expressions add optionality and choice operators to set of basic regex meta characters, as shown in Table 2.8. Table 2.9 shows examples for using these operators.

⁴¹Source: https://en.wikipedia.org/wiki/Regular_expression

RegEx	Matches
.at	"hat", "cat", "bat", "4at", etc.
[hc]at	"hat", "cat"
[^b]	all strings matched by .at except "bat"
[^bc]	all strings matched by .at except "bat" and "cat"
^[bc]at	"bat" and "cat" at start of text
[bc]at\$	"bat" and "cat" at end of text
\[.\]	any single character surrounded by [and], e.g. "[a]", "[7]", etc.
s.*	character "s" followed by zero or more characters, e.g. "s", "saw", "s3w96.7", etc.

Adapted from https://en.wikipedia.org/wiki/Regular_expression

Table 2.7: Basic Regular Expression Examples

Meta character	Description
?	Matches preceding element zero or one time
+	Matches preceding element one or more times
	Matches either the expression before or after the choice operator

Adapted from https://en.wikipedia.org/wiki/Regular_expression

Table 2.8: Extended Regular Expressions

Over the years, different types or "dialects" of regular expressions have been developed for or within different programming languages. One popular dialect is that used in the Perl programming language or the Vim text editor. One important way in which they differ is in the character classes they provide as shortcuts for specifying a set of characters to match. Table 2.10 shows an excerpt of the most frequently used character classes.

Regular expressions are a fundamental tool in text processing and manipulation, offering a robust and efficient method for pattern matching and string analysis. Their versatility makes them an essential skill in many programming and data-related tasks. Regular expressions are available in all programming languages and statistics and ana-

RegEx	Matches
[hc]?at	"at", "hat", "cat"
[hc]*at	"at", "hat", "cat", "chat", "chchchat", etc.
[hc]+at	"hat", "cat", "chat", "chchchat", etc.
cat dog	"cat" or "dog"

Adapted from https://en.wikipedia.org/wiki/Regular_expression

Table 2.9: Extended Regular Expression Examples

	Perl/Vim	ASCII	POSIX
Digits	\d	[0-9]	[[:digit:]]
Non-digits	\D	[^0-9]	
Lowercase letters	\l	[a-z]	[[:lower:]]
Uppercase letters	\u	[A-Z]	[[:upper:]]
Alphanumeric chars	\w	[A-Za-z0-9_]	
Non-word chars	\W	[^A-Za-z0-9_]	
Whitespace	\s	[\t\r\n\f]	[[:space:]]
Non-whitespace	\S	[^ \t\r\n\f]	

Table 2.10: Character classes in Regular Expressions

lytics software tools and allow a basic level of text processing and manipulation.

Hands-On Exercise

- Specify a RegEx to match Canadian postal codes:

<https://www.canadapost-postescanada.ca/cpc/en/support/articles/addressing-guidelines/postal-codes.page>

- Specify a RegEx to match a full RFC 3339 date with timezone, such as "2023-11-14T20:42:53-04:30"
- Challenge:* Specify a RegEx that matches any ISO 8601 date-time format

Levenshtein Distance

The Levenshtein distance is a metric of similarity of two text fragments. It is a type of string-edit distance, in that it measures the lowest number of *insertion*, *deletion* and *substitution* operations of individual characters to transform one text fragment into the other. The operations may be equally weighted or be differentially weighted, for example to penalize deletion operations more than insertion operations. The recursive definition is shown in Figure 2.11.

As an example, consider the two strings "kitten" and "sitting". The (unweighted) Levenshtein distance between the two is 3. In the first edit, the "k" is substituted with an "s", then the "e" is substituted with an "i" and finally a "g" is inserted at the end.

Hands-On Exercise

Determine the Levenshtein distances between the following:

- Last five digits of your student number and "12345"
- The words "Nunavut" and "Nunatsiavut"
- The words "Inuktitut" and "Innuttitut"
- The words "Mikak" and "Micock"

$$\text{lev}(a, b) = \begin{cases} |a| & \text{if } |b| = 0, \\ |b| & \text{if } |a| = 0, \\ \text{lev}(\text{tail}(a), \text{tail}(b)) & \text{if } \text{head}(a) = \text{head}(b), \\ 1 + \min \begin{cases} \text{lev}(\text{tail}(a), b) \\ \text{lev}(a, \text{tail}(b)) \\ \text{lev}(\text{tail}(a), \text{tail}(b)) \end{cases} & \text{otherwise} \end{cases}$$

https://en.wikipedia.org/wiki/Levenshtein_distance

Figure 2.11: Levenshtein Distance

Images

Image data is the representation of visual information in a digital format. There are two primary types of image formats: vector and raster.



https://commons.wikimedia.org/wiki/File:Persian_sand_CAT.jpg

images is dependent on their resolution. Scaling up a raster image can lead to a loss in quality, known as pixelation. Common formats are JPEG, and PNG, which use a lossy compression, that is, in reducing the file size, image detail may be lost. In contrast, the TIFF format uses lossless compression, retaining the full information of an image.

Image data in analytics is typically in a raster format using the RGB colorspace, which describes colours in terms of their red, green, and blue components⁴³. Hence, each pixel is described by 3 bytes (color components range from 0 to 255) and a full image can be thought of conceptually as a $3 \times X \times Y$ array of values between 0 and 255. For image analytics, images in a compressed format such as JPEG, PNG or TIFF must be decompressed to the full set of $X \times Y$ pixels, and the three RGB values are usually scaled to a range between 0 and 1.

Typical image analysis tasks include object detection and counting, object classification, image segmentation, and image retrieval. Object classification or image classification categorizes an entire image, or specific objects within an image, into predefined classes. This is commonly used for social media analysis or applications like photo tagging. Object detection involves identifying and locating objects within an image.

⁴²<https://www.w3.org/TR/SVG2/>

⁴³Another widely used colour space is CMYK, where a pixel's color is described in terms of its cyan, magenta, yellow, and black components.

This task goes beyond merely recognizing what objects are present; it also determines where they are in the image. Typically, object detection algorithms output a bounding box for each detected object, specifying its coordinates within the image. Object detection is widely used in applications such as surveillance, face detection, and autonomous vehicles. Image segmentation divides an image into multiple segments with the aim of simplifying or changing the representation of an image into a form that is more meaningful and easier to analyze. Image segmentation is used in medical imaging, machine vision, and object tracking. Image retrieval involves searching and retrieving images from a large database based on the content of the images themselves. It typically involves extracting features like color, texture, and shape from the images and using these features to find similar images in a database.

Business applications of image analysis include robotics, character and handwriting recognition in documents for process automation, security (identity verification, fraud detection, etc.) and manufacturing (defect detection, etc.).

Hands-On Exercise

1. Identify a specific business problem that can be addressed by analyzing image data
2. What image data would you need to address the problem?
3. What would you wish to do with the image data?
4. Where might you get this image data?

Video

Video data consists of a sequence of images ("frames") displayed at a certain rate (frame rate) to create the illusion of motion. Accompanying audio tracks are synchronized with these frames. Video data can be complex due to the need to balance quality, resolution, compression, and file size. Conceptually, video is a series of image *frames* in raster image format, i.e. a $T \times 3 \times X \times Y$ array of RGB values between 0...255 (where T refers to the set of frames over time).

However, in practice, video data is heavily compressed in video files as specified by different video formats. Each format has its compression techniques and algorithms, impacting the video's quality, size, and playback compatibility. A video *codec* (compressor-decompressor) is a software or hardware tool that compresses (encodes) and decompresses (decodes) digital video in a particular format to reduce file size and bandwidth requirements for storage or transmission. Popular video formats (codecs) are H.264, H.265, AVC, and AV1. A video *container format* is a file format that can contain various types of data, including video, audio, subtitles, and metadata. The container format determines how the data streams are organized and synchronized to each other. Popular container formats include MPEG-4, MKV, AVI, VOB and WebM.

Typical video analytics tasks include object detection, object recognition, object motion detection, object or background dynamic masking/blurring, event detection and

classification (errors, exceptions), and activity detection and classification. Object detection in video involves identifying and locating objects within a frame or series of frames. This task typically recognizes and tracks multiple objects over time, often in real-time. Object recognition goes a step beyond detection to classify the detected objects into predefined categories, such as identifying specific types of vehicles, animals, or other objects within a video. Motion detection involves identifying moving objects in the video. It is crucial in surveillance systems to detect unusual or suspicious movements or to track the movement of specific objects or people over time. Dynamic masking or blurring is used to obscure or protect portions of the video image, such as faces or license plates, to maintain privacy or comply with regulations. Event detection involves identifying specific events within a video, such as errors, exceptions, accidents, or other significant incidents. Classification categorizes these events into predefined types to facilitate appropriate responses or further analysis. Activity detection involves recognizing and categorizing the actions or behaviors of objects or people in the video, such as walking, running, or using machinery. This can be used in various settings, from analyzing customer behavior in retail to monitoring patient activities in healthcare.

Hands-On Exercise

1. Identify a specific business problem that can be addressed by analyzing video data
2. What video data would you need to address the problem?
3. What would you wish to do with the video data?
4. Where might you get this video data?

2.3 Metadata

Metadata is often described as "data about data." It provides information about, or documentation of, other data managed within an application or data store. Metadata is crucial for understanding, managing, and using the actual data effectively.

Metadata can describe authorship and ownership of the data, e.g. who created or owns it. It can also describe licensing and legal information, such as what one is allowed to do with a data set, what purposes it may be used for, whether it may be copied or redistributed, etc. Metadata can also provide information about when, where, and how data was collected or processed. It can specify the meaning of fields in tabular data, or of properties in graph databases. Metadata can be used to describe validation rules for data. Finally, metadata may be technical information, for example, describing the encoding or serialization format of the data.

Some data formats allow meta-data to be embedded within them, such as popular image or video formats. For other data formats, such as CSV files, metadata may be provided as a separate document or simply as a text file.

Hands-On Exercise

1. With your cell phone camera, take a selfie
2. Identify the meta-data that your phone camera embedded in this photo

2.4 Data Quality and Data Provenance

Data quality refers to the condition or fitness of data to serve its intended purpose in a given context. Poor data quality will lead to poor predictions, prescriptions, and decisions. Data quality has a long history both in research and practice⁴⁴. There are many dimensions to data quality, depending on the kind of data and the purpose for which the data is intended. Different authors and sources will list different dimensions; Table 2.11 gives an overview of the most important aspects of data quality.

Achieving, maintaining and ensuring data quality is a continuous process that involves regular monitoring, cleaning, standardization, and validation of data. Because of its importance, and the significant effort involved in it, data quality management is often a centrally located responsibility of the Chief Information Officer (CIO) or other senior management of an organization. Typically, organizations aim to have procedures and policies into place that govern data quality and how to achieve and maintain it.

Dimension	Example Considerations
Accuracy	Error rate for numerical data
Availability	Cost or ease of retrieval or collection or licensing
Completeness	Incomplete data may lead to bias
Conformity	Conforms to internal and/or external standards
Consistency	Free from internal contradictions
Integrity	Complies with validation rules, data types, and schema
Precision	Measurement precision of values
Relevance	Usefulness for purpose
Reliability	Consistency of repeated data points
Timeliness	Latency, currency, "age"
Traceability	Auditable provenance, verifiable source

Based in part on: Richard Y. Wang & Diane M. Strong (1996) Beyond Accuracy: What Data Quality Means to Data Consumers, Journal of Management Information Systems, 12:4, 5-33, DOI: 10.1080/07421222.1996.11518099

Table 2.11: Data Quality Dimensions

⁴⁴For a seminal academic reference, see Richard Y. Wang & Diane M. Strong (1996) Beyond Accuracy: What Data Quality Means to Data Consumers, Journal of Management Information Systems, 12:4, 5-33, DOI: 10.1080/07421222.1996.11518099

Data provenance refers to the documentation or tracing of the origins, lineage, and lifecycle of data. It encompasses recording information of the inputs, entities, systems, and processes that influence the data of interest, providing a record of the data and its origins. Data provenance is crucial for understanding the context, derivation, and rationale behind the data, making it an essential aspect of data management and an important prerequisite for data quality.

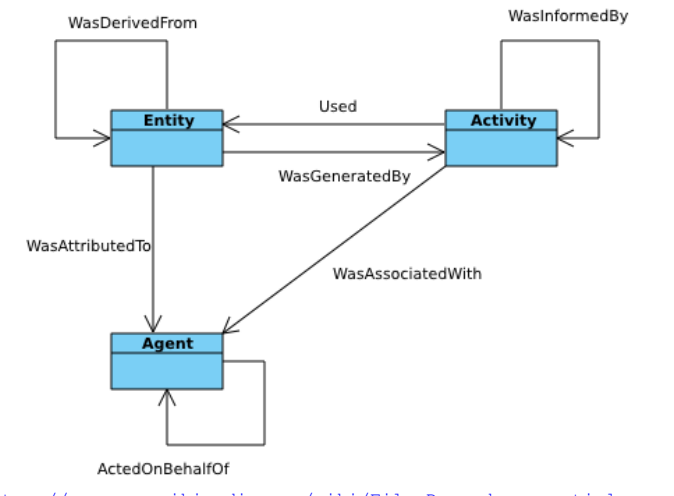
Source tracking identifies where the data comes from, including the original source of the data and any intermediate sources. Tracking data transformation or processing keeps a record of how the data has been altered, transformed, or processed from its original state. This includes changes in format, structure, or content. Tracking of ownership and responsibility documents who has handled or managed the data throughout its lifecycle. Versioning information keeps track of different versions or states of the data over time.

Provenance information helps in assessing the reliability and trustworthiness of data. Knowing the source and history of data can establish confidence in its accuracy and validity. Data provenance also provides transparency into the data's history, ensuring accountability for the data's quality and integrity. Understanding the provenance of data can aid in identifying when and where errors were introduced into the dataset. This facilitates more effective error correction and data cleansing.

Figure 2.12 shows a recommendation by the World Wide Web Consortium (W3C) of the basic elements of a framework to maintain data provenance records. Agents are associated with activities that use or create data entities. In turn, data entities are derived from other data entities, and are attributed to agents, e.g. as creators. Figure 2.13 shows an example diagram of a provenance record using this framework. The figure shows agents playing the roles of contributor and editor with respect to an editing activity of a data object that was generated by the editing activity.

Data provenance is about asking and answering questions related to the data and all that happened to it. Important questions include:

- How was the data collected? What errors could have occurred?
- Who collected the data? Is it a trustworthy source?
- When were the data collected? Are they still valid?
- Are all the data collected? Are the data biased?
- Can the data collection be verified/audited/repeated?
- How was the data processed? What mistakes could have been made?
- Was anything omitted or added?
- Who processed the data? Is it a trustworthy party?
- Can the processing be verified/audited/repeated?
- What do different data fields mean?



https://commons.wikimedia.org/wiki/File:Prov_dm-essentials.png

Figure 2.12: Data Provenance Framework Basics

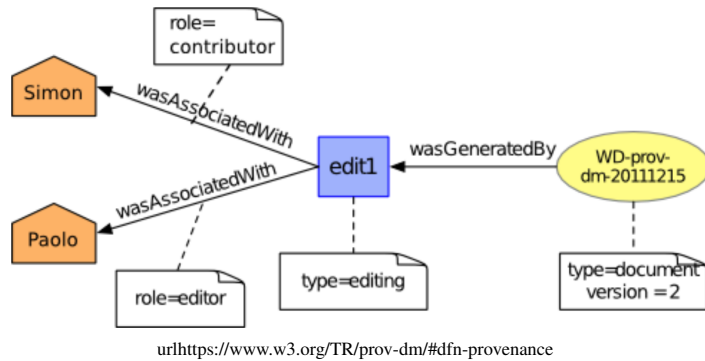


Figure 2.13: Data Provenance Framework Example

- What are the units of measurement?
- What is the level of aggregation?
- Were data sources combined? Are the different sources consistent with each other and of the same quality?
- Are the data accurate? How high are the error rates and the levels of precision?
- Can the data be validated? What are the validation rules for the data? Was the data validated?
- How can errors be detected and/or corrected?
- Are the data usable in a technical and legal way?

2.5 Data Cleaning and Validation

Data cleaning is a critical step in the data analysis process and contributes to high quality. It involves the identification of errors and inconsistencies in the data and their correction. Data correction can mean different things in different situations, from simply omitting erroneous data, to "clipping" numerical data within certain ranges, standardizing or normalizing textual data (for example all lowercase, word stemming, etc.), imputing missing data (for example, by using the mean or some more sophisticated method), etc. The ultimate aim is improve the data quality and therefore the quality of the analysis results themselves. Data cleaning typically involves a number of steps:

1. Auditing

This step identifies anomalies and inconsistencies. It requires a thorough understanding not only of the data but also how the data was collected, and what the data is intended to describe or represent, that is, the domain. Only then can errors be identified (e.g. based on plausibles mistakes during data collection) and the internal consistency of the data (e.g. based on what is plausible in the domain) be evaluated.

2. Validation

Ensure data conforms to rules and constraints. This requires first identifying any rules for data coding or data consistency constraints that should apply. Next, data that violates these rules and constraints can be identified. Example of data validation rules are:

- Encoding or serialization rules, e.g. with Regex
 - *Example:* Are all phone numbers of the format

$$\stackrel{\wedge}{(}[0-9]\{3\})[-]?[0-9]\{3\}[-]?[0-9]\{4\}\$$$
- Data type constraints
 - *Example:* Are all sales prices numbers?
- Range constraints
 - *Examples:* Are prices > 0? Are sales numbers < 1000?
- Cross-field validation
 - *Example:* If province is NL, then phone area code must be 709 or 879

3. Cleaning

Cleaning involves the transformation and correction of data, including identifying how to deal with missing values. This may also include bringing data to standardized formats, e.g. transforming numbers, dates, standardizing abbreviations and spelling, etc. Numerical data may be clipped or constrained to certain ranges, and inconsistencies between different data items must be resolved.

- *Data Transformation:* Convert data into required format or structure. For example,
 - One row for each observation, case, or event
 - Create case or event identifiers
 - *Data Imputation:* Replacing missing values with estimated or default values, or removing missing values entirely. Be mindful that:
 - Missing values may have different meanings
 - Data removal may bias data
 - Estimating values may introduce errors
 - *Data Correction:* Correct or remove erroneous data
 - Importantly, data correction requires access to correct data, which may not be available or must be provided by other, secondary sources.
4. *Duplicate Removal:* Ensure uniqueness of data. Duplicates may be real duplicates or simply the result of different spelling or abbreviations or other data entry mistakes. In any case, duplicates can bias analysis results that rely on sums, counts, variances or other statistics. Consider the following example of names:
- *Example:* Rebekah Uqi Williams (Commissioner of Nunavut (2020–2021))
 - *Abbreviations:* Rebekah U. Williams; Rebekah Williams, R.U. Williams
 - *Order:* Williams, Rebekah Uqi; Williams, Rebekah U.; Williams, R., ...
 - *Spelling:* Rebekah; Rebecca; Rebeccah; Rebeckah; Rebecka, ...
 - *Misspellings:* Reebkah, Rebkah, Wililams, Willaims, ...
5. *Harmonization:* Merge datasets from different sources and ensure consistent formats and scales. For example, standardize date and number formats, standards units of measure, etc.
6. *Standardization:* Bring data into a standard format. Chapter 2 showed that standards exist in many areas for many data types. It is important for further analysis to ensure data complies with standards, to be able to easily, efficiently and effectively use tools for further analysis.
7. *Quality Assessment:* Ensure cleaning has been effective. Re-assess the resulting data set on data quality aspects.

Cleaning, transformation, and correction of data is *subjective* and requires a domain or business expert with *expert knowledge* of the data and its provenance, the metadata, the validation rules, and the application domain.

In practice, cleaning, transformation, and correction of data takes approximately 80% of a data analyst's time, while actual analysis takes only 20% of their time. This is sometimes called "data wrangling".

2.6 Data Sources

Data used for business analytics can be internal to an organization or acquired from external sources. Often, the data required to address a particular analytics problem is a combination of internal and external data, i.e. the internal data is *enriched* with external data.

Internal data sources may be operational computer systems, such as the HR, payroll, accounting, logistics, manufacturing, sales systems and many others. These systems provide operational data about human resources, finances, goods movements, etc. Another source of data are *data-rich* products. Since the 2010s, companies are increasingly selling products that include a variety of sensors, with the ability for the sold products to provide information back to the manufacturer or some other organization. Such data-rich products, whether they are cars⁴⁵ or teddy bears⁴⁶, can provide a vast amount of rich data relating to the operation of the device and the customer that uses the device. Most computer systems also keep technical logs. The most prominent example are web-server logs, but many other computer systems do as well. Such logs provide information about who accesses what information or performs what operation at what time. Rich information can also be obtained from message data, whether those messages are emails that pass through the company's email servers, customer service chat interactions, or call center audio recordings. Finally, data may be directly collected from humans for a specific project or purpose, for example in the form of employee or customer surveys.

External data may be public or private. The increasing popularity and use of analytics has fuelled the publication of many data sets by governments, international institutions, and companies interested in furthering analytics applications and insights. These data sets are now easier to access than ever. Table 2.12 provides examples of public, external data sources and where to find them. However, when using these data sets it is important to critically assess their provenance and quality.

If the required data is neither internally nor publicly available, there exist many sources to purchase data, especially financial market and consumer data. Such data may come directly from services companies or may be provided by data brokers who aggregate data from a variety of sources to increase the value of the data. Table 2.13 shows some examples of private external data sources.

Data licenses provide the legal framework and specify permissions governing the use,

⁴⁵ <https://foundation.mozilla.org/en/privacynotincluded/articles/its-official-cars-are-the-worst-product-category-we-have-ever-reviewed-for-privacy/>

⁴⁶ <https://arstechnica.com/information-technology/2017/02/creepy-iot-teddy-bear-leaks-2-million-parents-and-kids-voice-messages/>

Government Agencies	
Statistics Canada	https://www.statcan.gc.ca/en/start
Open Government Canada	https://search.open.canada.ca/opendata/
US Census Bureau	https://www.census.gov/
US Bureau of Labor Statistics	https://www.bls.gov/
International Institutions	
OECD	https://data.oecd.org/
Worldbank	https://data.worldbank.org/
EU	https://data.europa.eu/en
WHO	https://www.who.int/data
Data Set Search Engines	
Google Dataset Search	https://datasetsearch.research.google.com/
GitHub Data Set Search	https://github.com/search?q=datasets&type=repositories
Social Media Companies	
X	https://help.twitter.com/en/rules-and-policies/x-api
Google	https://developers.google.com/gdata
Facebook/Meta	https://developers.facebook.com/docs/graph-api/overview/
ML/AI Project Communities	
Kaggle	https://www.kaggle.com/
HuggingFace	https://huggingface.co/datasets
Google Cloud	https://console.cloud.google.com/marketplace/browse
Google Research	https://research.google/resources/datasets/
AWS Data Sets	https://registry.opendata.aws/
Azure Data Sets	https://azure.microsoft.com/en-ca/products/open-datasets

Table 2.12: Examples of Public External Data Sources

Services Companies	
Financial services institutions	e.g. Bloomberg https://www.bloomberg.com/professional/product/data/
Telecommunications providers	e.g. Telus Insights https://www.telus.com/en/business/medium-large/enterprise-solutions/big-data-analytics
Mobile applications	e.g. The Weather Network https://www.pelmorex.com/en/data/
Data Brokers	
LiveRamp (formerly Axiom)	http://www.liveramp.com/
Experian	https://www.experian.com/
CoreLogic	http://corelogic.com/
Nielsen	http://nielsen.com/
DataAxeCanada (formerly InfoCanada)	https://www.dataaxlecanada.ca/

Table 2.13: Examples of Private External Data Sources

redistribution, and modification of data. They dictate how data can be shared and used, outlining the rights and restrictions placed on the data by its owner or creator. Data

licensing is especially important in the era of big data and open data initiatives, where data is often shared and reused across various domains and applications.

The creator of data is typically the owner and obtains copyright to the data. Copyright impacts how the data can be legally used and shared. Usage rights or licenses specify what others can and cannot do with the data. Licenses may include permissions for using the data. Licenses also govern whether the data can be redistributed to third parties and under what conditions. They also dictate whether the data can be modified and how derivative works (new creations based on the original data) are to be handled. Importantly, data licenses may also specify whether the data can be used for commercial purposes at all, and any conditions or restrictions on access to the data.

Different kinds of licenses exist for data. Open data licenses allow data to be freely used, modified, and shared by anyone. Examples include the Creative Commons licenses⁴⁷ and the Open Data Commons licenses⁴⁸. Additionally, many governments and international institutions provide data with open licenses. In contrast, proprietary licenses restrict usage to certain conditions set by the owner, which can include the requirement for payment, restrictions on redistribution, or limitations on the type of use (e.g., non-commercial only). Some data may be in the public domain and is not protected by copyright so that it can be freely used by anyone without restrictions.

Data licensing is a critical aspect of data governance and data management, particularly in contexts where data is shared and reused extensively. Understanding and complying with data licenses is essential for anyone involved in data analysis, software development, and research. Never assume permission to use the data for any particular purpose is given simply because the data is accessible. Check the meta-data of the data, or the web-site where the data is available. Some organizations may provide permission to use data or licenses upon request, while others may require the purchase of a license, especially if the data is to be used for commercial purposes.

⁴⁷<https://creativecommons.org/>

⁴⁸<https://opendatacommons.org/>

Hands-On Exercise

1. Identify data on the consumer price index (excluding living and transportation expenses) for Newfoundland & Labrador for the last 10 years
 - How was it collected? By who? When?
 - How was it processed? By who? What was done to it?
 - Is there meta-data available for it?
 - How do you assess the quality of the data on the data quality dimensions?
 - Under what license is it available to you to use?
2. Identify some IoT devices or sensors in your household
 - What information can they measure?
 - How and when is the information being collected? By who?
 - How could the information be erroneous or biased?
 - How would you assess the quality of the data?

2.7 Review Questions

Data Types

1. What is the equivalent of R's `numeric` data type in Python?
2. Explain the difference between R's `integer` and Python's `int` data type.
3. What term does the R statistical system use to indicate a missing value?
4. How does Python represent a missing value?
5. What is the term used by SQL to denote a missing value?
6. If you are working with both Python and R, what considerations should you keep in mind regarding data types when transferring data between these two languages?
7. Give an example where a data type in R might not have a direct equivalent in Python or SQL.
8. Discuss the potential issues that might arise when working with missing values in data analysis.
9. Provide an example of a scenario where the meaning of a missing value can be ambiguous. How might this ambiguity impact data analysis?
10. In a dataset, you find that some entries are marked as `NA` in R, `None` in Python, and `Null` in SQL. How would you interpret these values?
11. What steps could you take to handle missing values before performing any statistical analysis?

Number Formats

12. Describe how integer numbers are represented in binary form. What does the first bit indicate?
13. How many bytes does a `float` (single precision number) occupy? Break down its composition in terms of sign, exponent, and fraction.

14. Discuss some of the complexities involved in writing out decimal numbers as plain text.
15. How do decimal point representations differ in various locales?
16. Explain how negative numbers and scientific notation are represented differently in various contexts.
17. Why is it important to verify the number format in a dataset before using it with a business analytics tool?

Text Format

18. Explain the purpose of the Unicode system and how it addresses the limitations of earlier encoding systems like ASCII.
19. Discuss the variety of characters, symbols, and scripts included in the latest version of Unicode (v15.1).
20. How does UTF-8 ensure backward compatibility with ASCII?
21. Why is it important for a business analyst to be aware of Unicode and its different encodings?
22. Discuss the implications of using Unicode-aware data storage, management, analytics, and visualization tools in a business setting.

Date Formats

23. Explain the challenges in handling different time zones in a global context.
24. How do ISO 8601 and RFC 3339 standards differ from each other?
25. Describe the variety of formats and separators used in writing dates across different locales.
26. Explain the challenges involved in performing arithmetic operations with years, months, and days due to their different lengths and conventions.
27. Discuss the implications of not universally accepting standards like ISO 8601 and RFC 3339 in data management and analytics.

Collection Types

28. Define what a collection data type is and explain how it differs from primitive data types.
29. Describe the characteristics of a list in Python and compare it with the list in R.
30. Discuss the structure and usage of dictionaries in Python.
31. Describe the properties of a vector in R and how it differs from a list.
32. Discuss the significance of mutable and immutable data types, providing examples from Python and R.

Tabular Formats

33. Explain the CSV (Comma-Separated Values) file format. What are its key characteristics?
34. According to RFC 4180, what are the standard conventions for formatting a CSV file?

35. Discuss the common variations and deviations you might encounter in CSV files that do not strictly adhere to the RFC 4180 standard.
36. How are line breaks typically represented in CSV files, and what are the common variations?
37. What challenges might you face when working with CSV files that do not conform to standards, and how could you address these challenges?

Document Formats

38. Define JSON and XML and explain their primary purpose in data interchange.
39. What are the commonalities and differences between JSON and XML? When would you prefer one over the other?
40. Describe the structure of a JSON document in terms of key-value pairs.
41. What types of values can be stored in a JSON document?
42. Explain how objects are represented in JSON. What delimits an object?
43. Describe how arrays are represented in JSON and how they differ from objects.
44. How can an array be nested within a JSON object, and vice versa?
45. What is the purpose of a namespace in XML documents? Provide an example.
46. How do elements and attributes differ in XML? In which situation would you choose an element? In which situation would you choose an attribute?

Text Data

47. What is text analysis and why is it important in extracting information from unstructured text data?
48. Describe named entity recognition and its application in content classification and data extraction.
49. Explain co-reference analysis and provide an example of how it helps in understanding text.
50. Discuss event and relationship extraction in text analysis, providing an example.
51. Describe sentiment analysis and its significance in areas like social media monitoring and market research.
52. What is document clustering and how is it used to organize and categorize text data?
53. Compare and contrast the symbolic, statistical, and deep learning approaches in text mining.
54. Discuss the impact of deep learning and neural networks on the field of text analytics, specifically mentioning models like RNNs, LSTMs, and transformers.

Regular Expressions

55. Define regular expressions and explain their primary purpose in text processing.
56. What are meta characters in the context of regular expressions?
57. Give examples of basic meta characters in regular expressions and explain their functions.
58. Describe the additional capabilities provided by extended regular expressions.

59. How do regular expressions differ from literal text searching?
60. Provide an example of a regular expression pattern and explain what it matches.
61. Discuss the challenges or limitations associated with using regular expressions.

Image Data

62. What are the two primary types of image formats? Describe the main characteristics of each.
63. Explain the concept of vector images. What are some common formats of vector images?
64. How do vector images maintain quality when scaled to different sizes? Provide a brief explanation.
65. Define raster images and explain how they are structured.
66. What is the main limitation of scaling raster images, and why does this limitation occur?
67. List some common raster image formats and mention whether they use lossy or lossless compression.
68. How are pixels represented in a typical raster image in terms of RGB values?
69. What are some common tasks involved in image analysis? Briefly describe each task.
70. Discuss the application of image analysis in business, providing at least three examples.

Video Data

71. What does a video frame represent in terms of data structure? Explain the notation $T \times 3 \times X \times Y$ in this context.
72. Discuss the role and importance of compression in video data.
73. What is a video codec? Give examples of popular video codecs and their general applications.
74. Explain the purpose of a video container format and list some common container formats.
75. Describe the concept of object detection in video analytics and how it differs from object recognition.
76. What is motion detection in video and why is it important in surveillance systems?
77. Discuss the purpose and applications of dynamic masking or blurring in videos.
78. Define event detection in video analytics and give examples of events that might be detected.
79. Explain activity detection in video and its potential applications in various industries.

Metadata

80. Define metadata and explain its significance in data management and utilization.
81. Describe how metadata can provide information about authorship and ownership of data.

82. Explain how metadata can be used to detail the collection and processing of data.
83. Give examples of technical information that metadata might describe.

Data Quality

84. Define data quality and explain why it is important in data management.
85. List and describe the various dimensions of data quality.
86. Discuss the processes involved in maintaining high data quality.
87. How does poor data quality affect predictions, prescriptions, and decisions?
88. Explain the concept of data cleansing and its role in data quality management.

Data Provenance

89. Define data provenance and its significance in the context of data management.
90. What types of information are typically included in data provenance records?
91. Explain how data provenance contributes to the reliability and trustworthiness of data.
92. Discuss the role of data provenance in error detection and correction in datasets.
93. Provide examples of questions that are important to ask when evaluating data provenance.

Data Sources

94. What are some examples of data-rich products, and what kind of data can they provide?
95. Discuss the role of web-server logs and other technical logs as sources of data.
96. What are the typical ways data is directly collected from humans for business purposes?

Data Licensing

97. Define data licenses and their importance in the context of data management.
98. Distinguish between open data licenses and proprietary licenses.
99. Explain the steps one should take to ensure compliance with data licenses.

2.8 Hands-On Exercises

Number Formats

1. Convert the decimal number 25 to its binary equivalent. Indicate the sign bit.
2. Convert the binary number 1101011 to its decimal equivalent.
3. Write the number 1234567.89 in four different formats, considering decimal points, digit grouping, and negative number representation.
4. Convert $5.12e3$ and $-3.04e-2$ to their regular decimal forms in two different locale styles.

5. Given a dataset with numbers in European format (comma as decimal separator), write a pseudo-code to convert them to the American format (dot as decimal separator).
6. Create a small program in a language of your choice to detect and convert scientific notation to standard decimal notation.

Character Formats

7. Use an online Unicode character table (like <https://www.unicode.org/charts/>) to find the Unicode characters for the letters in your name in a non-Latin script (e.g., Cyrillic, Greek, Arabic). Write the Unicode code points for these characters in both hexadecimal and decimal formats.
8. Choose a word in a language that uses non-ASCII characters. Find the Unicode code points for each character of the word. Convert these code points into UTF-8 encoded byte sequences. You can use online tools or write a simple program to do this.

Date Formats

9. Research a non-Gregorian calendar system (e.g., Hebrew, Islamic, or Chinese calendar). Convert today's date from the Gregorian calendar to your chosen calendar system. Discuss the key differences and similarities between the two calendar systems.
10. Choose three cities in different time zones. Convert 12:00 PM in your local time to the time in each of these cities. Discuss how time zone differences impact global communication and business.
11. Write a program or script to determine if a given year is a leap year in the Gregorian calendar. Test your program with a set of years, including at least one century year.
12. Write the current date and time in the formats specified by both ISO 8601 and RFC 3339. Discuss why such standardizations are important in data management and international communications.
13. Calculate the number of days between your birth date and today using a date arithmetic tool or programming library. Discuss the challenges you might face when calculating durations involving months and years due to their varying lengths.

Structured Data

14. Create a list with different data types, append a new element, and modify an element. Perform similar operations with a list in R.
15. Create a tuple in Python, attempt to modify it, convert it to a list, modify the list, and then convert it back to a tuple.
16. Create a dictionary in Python, add, modify, and retrieve values from it.
17. Create numeric and character vectors in R and apply various functions to them.
18. Convert a character vector in R into a factor and reorder its levels.

19. Create a matrix in R, access its elements, and perform matrix multiplication.
20. Convert one structured data type into another in both Python and R.

Tabular Formats

21. Manually create a CSV file using a text editor. Include a header row and at least 4 rows of data. Ensure that your CSV adheres to the RFC 4180 standard.
22. Write a simple program or script in a language of your choice (like Python or R) to read the CSV file you created and print out each row. Handle potential errors like missing fields or incorrect formatting.
23. Modify your CSV file to include a non-standard delimiter (like a semicolon) and mixed quotes. Adjust your program or script to correctly parse this modified CSV file.
24. Extend your CSV file by adding a column that includes complex data types (like lists or sets). Modify your parsing program to correctly interpret and display these complex data types.

Document Formats

25. Create a JSON object that represents a book, including properties such as title, author, publication year, and genre. Validate the JSON object using an online JSON validator.
26. Extend the book JSON object to include a nested object for the author, with properties like name, birth year, and nationality. Validate and format the updated JSON object.
27. Create a JSON array representing a book series, containing several book objects. Validate the JSON array to ensure proper formatting.
28. Take a simple dataset (e.g., a CSV file with student records) and convert it to a JSON format. Validate the converted JSON data.

Regular Expressions

29. Write a regular expression to match email addresses in a text. Test your expression on a set of sample strings to check its accuracy.
30. Create a regular expression using meta characters to match any date in the format "dd/mm/yyyy". Validate your RegEx with various date strings.
31. Write a RegEx to find all the hyperlinks (URLs) in a given HTML document.
32. Develop a RegEx to identify phone numbers in different formats (e.g., 123-456-7890, (123) 456-7890). Test the RegEx for various phone number formats to ensure its versatility.
33. Choose a programming language and use its RegEx library to split a paragraph into sentences. Ensure that the RegEx correctly handles periods used in abbreviations.

Levenshtein Distance

34. Determine the Levenshtein distance between "intention" and "execution".

35. What is the Levenshtein distance between a string and an empty string? Verify your answer using the strings "algorithm" and "".
36. Compute the distance between two identical strings, such as "database" and "database".
37. Given a list of words, ["apple", "apply", "apology", "propel"], find the word with the smallest Levenshtein distance to "aply".

Data Sources

38. Identify various internal data sources within a hypothetical or real organization (e.g., sales, HR). Discuss the types of data each source provides and its potential use in analytics.
39. Research and compare two different data licenses (e.g., a Creative Commons license and a proprietary license). Summarize the key permissions and restrictions of each license. Discuss the potential implications of these licenses on data usage in a business context.
40. Study a real case where data licensing played a critical role in a project or product. Identify the licensing issues that were involved and how they were addressed. Reflect on the lessons learned and how they apply to data management practices.

Chapter 3

Managing Tabular Data with Relational Databases

3.1 Introduction

The relational database model, developed by Edgar F. Codd in 1970, is a fundamental approach in data organization and management. It structures data in tables, or relations, comprising rows and columns, where each row signifies a record, and each column denotes a field within the record. This model is grounded in principles like tables, primary keys for unique record identification, foreign keys for inter-table relationships, and data integrity through constraints. The Structured Query Language (SQL) significantly improved the usability of data management with relational databases.

The 1970s marked the theoretical development of the relational model, focusing on data independence and efficient access. The 1980s witnessed its commercialization with the advent of relational database management systems (RDBMS) such as Oracle, IBM DB2, and Microsoft SQL Server, which became staples in enterprise applications. The 1990s saw the internet's rise bring scalability and distribution challenges to the forefront, leading to the popularity of open-source RDBMS like PostgreSQL and MySQL.

In the 2000s, with the onset of Big Data and the advent of NoSQL databases, the relational model faced new challenges. However, it continued to evolve, adapting features to handle non-relational data and integrating with cloud services. Relational databases have maintained their relevance and are extensively used in various sectors, including cloud computing, mobile applications, and big data analytics. The relational database model's focus on simplicity, flexibility, and accuracy has solidified its standing as a cornerstone in data management.

The relational database model offers several benefits and advantages, making it a popular choice for a variety of data management needs. One of its primary strengths is the

simplicity of its design, which organizes data into tables, making it intuitive and easy to understand. This tabular structure facilitates efficient data retrieval and manipulation, especially with the use of SQL, a powerful and standardized query language that enhances the accessibility and handling of data.

Another significant advantage is data integrity. The relational model enforces rules through primary and foreign keys, ensuring that relationships between data are logically maintained and that the data remains consistent and accurate. This is crucial for applications where data reliability is paramount.

The model's flexibility is also a key benefit. It can easily accommodate changes in the database structure without disrupting the existing data. This adaptability makes it suitable for a wide range of applications, from small-scale projects to large, complex enterprise systems.

Moreover, relational databases support ACID (Atomicity, Consistency, Isolation, Durability) properties of transactions (that is, updates to the data), guaranteeing reliable transaction processing and robust data management, especially in multi-user environments. This ensures that even in the event of system failures or concurrent data access, the integrity of the data is maintained.

The relational model's widespread adoption has led to a rich ecosystem of tools and technologies, providing users with extensive support and resources. This includes advanced features like indexing, which enhances performance, and comprehensive security measures for data protection.

3.2 Constraints and Data Types

In relational database management systems (RDBMS), constraints are essential for ensuring the integrity of the data. Constraints can be categorized into two main types: column constraints and table constraints. Additionally, each table column is of a certain primitive data type, allowing only certain types of values to be inserted. Constraints and typing ensure data quality, in that data conforms to expected rules. Table 3.1 shows an overview over commonly used datatypes.

Columns constraints are rules that are applied to individual columns, while table constraints apply to combinations of columns or the entire table. Many constraints can be specified both for a single column as well as a combination of columns.

- The *NOT NULL* constraint can only be applied to individual columns and prevents NULL values from being entered into a column, ensuring that every record has a value for that column.
- The *UNIQUE* constraint ensures that all values in a column are distinct, preventing duplicate entries. The UNIQUE constraint can also be used at the table level to ensure that a specific combination of values across different columns is unique for all records in the table.

Name	Description
<code>bigint</code>	signed eight-byte integer
<code>bit varying (varbit)</code>	variable-length bit string
<code>boolean</code>	logical Boolean (true/false)
<code>character varying (varchar)</code>	variable-length character string
<code>date</code>	calendar date (year, month, day)
<code>double precision (float8)</code>	double precision floating-point number (8 bytes)
<code>integer (int, int4)</code>	signed four-byte integer
<code>interval</code>	time span
<code>json</code>	textual JSON data
<code>jsonb</code>	binary JSON data, decomposed
<code>money</code>	currency amount
<code>numeric (decimal)</code>	exact numeric of selectable precision
<code>real (float4)</code>	single precision floating-point number (4 bytes)
<code>smallint (int2)</code>	signed two-byte integer
<code>text</code>	variable-length character string
<code>time</code>	time of day (no time zone)
<code>time with time zone (timetz)</code>	time of day, including time zone
<code>timestamp</code>	date and time (no time zone)
<code>timestamp with time zone, (timestamptz)</code>	date and time, including time zone

(Source: <https://www.postgresql.org/docs/current/datatype.html>)

Table 3.1: Primitive Data Types in SQL and PostgreSQL with aliases in parentheses. *Emphasized* entries are not contained in the SQL standard, they are PostgreSQL extensions.

- The *CHECK* constraint can be applied at the column level or at the table level. The *CHECK* constraint allows specifying a condition that each value in a column must satisfy. At the table level, it allows for more complex conditions that involve multiple columns.
- The *PRIMARY KEY* constraint is a combination of NOT NULL and UNIQUE, uniquely identifying each record in a table. The *PRIMARY KEY* constraints can also be applied at the table level to specify that a combination of columns uniquely identifies each record.
- The *FOREIGN KEY* constraint is used to link columns in different tables, establishing a relationship between them. It ensures that values in a column or combination of columns must exist in the referenced column or combination of

CREATE TABLE	Create a new table with specified columns and constraints
DROP TABLE	Deletes a table and all its contents
INSERT	Inserts a row of data values into a table
UPDATE	Updates/modifies data values in a table
SELECT	Retrieves data values from one or more tables

Table 3.2: Basic SQL Commands

columns. The referenced columns may be in the same table, so that the constraint expresses a *unary* relationship, or in another table, so that the constraint expresses a *binary* relationship. Together with NOT NULL constraints, this allows the representation of optional or mandatory relationships.

3.3 Introduction to SQL and PostgreSQL

Despite its name, SQL serves as a language not only for querying but for data definition, data manipulation, data access control, transaction control, and querying. SQL has been standardized by the American National Standards Institute (ANSI) and the International Organization for Standardization (ISO), ensuring a consistent syntax and set of features across different database systems. However, many database systems extend standard SQL with proprietary extensions to enhance functionality and performance. Despite these variations, the core elements of SQL remain widely consistent, contributing to its status as the lingua franca of database management.

This section covers only the most basic aspects of SQL, insofar as they are necessary to understand the relational database schema and to use SQL to query data for descriptive data analytics. The most important SQL commands are listed in Table 3.2. For more further information, consult the relevant sections of the PostgreSQL documentation on data definition¹, data manipulation², data queries³ and primitive data types⁴.



The PostgreSQL RDBMS (relational database management system) is installed in the course virtual machine or can be downloaded from the PostgreSQL website⁵. A DBMS is a background computer application without a user interface. It is typically used by other computer applications, such as accounting software to store financial information, a logistics management software to store information about shipments, a customer relationship management system to store information about customers and marketing campaigns, etc.

End users can interact with a DBMS using administration software, such as the basic

¹<https://www.postgresql.org/docs/current/ddl.html>

²<https://www.postgresql.org/docs/current/dml.html>

³<https://www.postgresql.org/docs/current/queries.html>

⁴<https://www.postgresql.org/docs/current/datatype.html>

⁵<https://www.postgresql.org/download/>

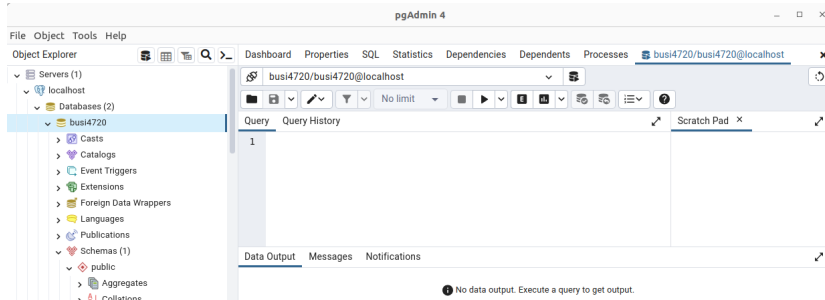


Figure 3.1: pgAdmin Query tool

”psql” command line software or a graphical application like ”pgAdmin” or ”DBeaver”. The desktop version of pgAdmin and DBeaver are installed in the course virtual machine, or can be downloaded from their websites⁶. They provide easy-to-use tools for creating tables and querying data, but this section focuses on using the SQL language instead.

A DBMS runs on a single computer (”server”) or, if the amount of data is very large, distributes the data across a cluster of multiple computers. Different DBMS differ in their performance, the ease with which data can be distributed, and the scalability to very large clusters. However, from the users perspective, these technical considerations are largely invisible. *When connecting to a DBMS that runs on your own computer, use the computer name ”localhost”.*

A DBMS can manage multiple databases. A *database named ”busi4720” has already been created in your course virtual machine, using the CREATE DATABASE command.* pgAdmin and DBeaver also have the ability to show the SQL command that creates every element in a DBMS, including databases, tables, and constraints. This is useful to understand exactly what elements are contained in a database or in a table and any constraints imposed upon them.

Every database can have multiple schema. A schema is a collection of tables with their columns and constraints, as well as related elements such as functions, procedures, triggers, views and others. In PostgreSQL, every database contains the schema ”public”. This is the default schema and is used when no other schema is specified.

When the pgAdmin application is initially launched, it will connect to the DBMS that is running on the local machine (”localhost”) with the username ”busi4720” (its password is ”busi4720”) and will show an ”Object Explorer” in the left part of the application. This allows navigation and exploration of the contents of this DBMS, as shown in Figure 3.1.

Similarly, when the DBeaver application is first started, it will also connect to the DBMS that is running on the local machine and will show a ”Database Navigator” in

⁶<https://www.pgadmin.org/download/>, <https://dbeaver.io/download/>

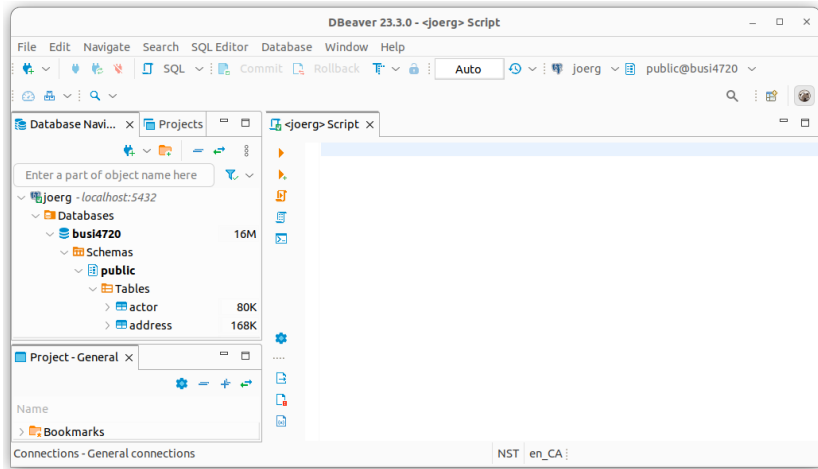


Figure 3.2: DBeaver database tool

```

busi4720@busi4720vm:~$ psql
psql (16.0 (Ubuntu 16.0-1.pgdg22.04+1))
Type "help" for help.

busi4720=# \conninfo
You are connected to database "busi4720" as user "busi4720" via socket at port "5432".
busi4720=#

```

Figure 3.3: psql command line tool

the left part of the application window (Figure 3.2). Navigate the contents of the DBMS to the "busi4720" database and the "public" schema.

The basic "psql" command line tool can be started by typing `psql` into a terminal window. The course virtual machine is configured to provide automatic access to the "busi4720" database. The database connection can be confirmed by executing the `\conninfo` command in psql. Use psql options to specify other connections using the template `psql -d dbname -h hostname -u username`. Figure 3.3 shows the psql command line tool in a terminal window.

The examples and exercises in the remainder of this chapter refer to the "busi4720" database.

3.4 Data Definition in SQL

Tip: SQL commands are traditionally written in upper case letters and this is done here as well. However, SQL is not case sensitive, so that capitalization does not actually matter. Traditionally, an SQL command must end with a semicolon. This is done here as well, although some DBMS may no longer require this.

For this example, assume that your database will be used to store information about products. The CREATE TABLE data definition command in SQL is used to create tables, their columns, and constraints. This first example creates a simple table with three columns to store product data.

Enter the SQL commands in the code block below.

- In psql, press **RETURN** to execute
- In DBeaver, press **CTRL-RETURN** to execute
- In pgAdmin, press **F5** to execute.

```
CREATE TABLE products (
    pcode integer,
    name  varchar(100) NOT NULL,
    price float4,
    PRIMARY KEY (pcode)
);
```

The table contains a column named "pcode" (to store the product code) that is of integer type, that is, it can contain whole numbers only. The table has a column called "name" of characters with a varying length and a maximum length of 100 characters to store product names. Additionally, a column NOT NULL constraint has been defined for this column, ensuring that a name always exists for a product. The table has a column named "price" that is of a single precision floating point type (4 bytes), that is, it can hold decimal point numbers to store product prices. The final line of the SQL command created a primary key constraint on the single column "pcode" to ensure that the product code must not be NULL and must be unique, ensuring that each row in the table represents a distinct product. Note that column "price" may be NULL, that is, may not contain values, because no NOT NULL constraints have been specified. This may be useful for example when the price has not been decided on or will be calculated later.

The following SQL code block creates a table for suppliers. The table has a similar structure to the products table and similar NOT NULL and PRIMARY KEY constraints.

```
CREATE TABLE suppliers (
    scode integer,
    name varchar(100) NOT NULL,
    city varchar(100),
    PRIMARY KEY (scode)
);
```

After creating tables with the data definition part of the SQL language, the data manipulation commands of SQL can be used to insert or update values in the tables. The following SQL code block inserts two rows into each of the tables that were just created. Enter the following SQL commands in the Query tool:

```
INSERT INTO products VALUES (1, 'Hex Bolt', 1.99);
INSERT INTO products VALUES (2, 'Round Bolt', 2.99);

INSERT INTO suppliers VALUES (1, 'Bolts Inc', 'HVGB');
INSERT INTO suppliers VALUES (2, 'Hardware Co', 'Cartwright');
```

The values for each row must be specified in the order in which the columns of the table are defined. For the "products" table this is first the product code, followed by the name, and finally the price. For the "suppliers" table, this is first the supplier code, then the name and then the city. There are many different variations on the basic INSERT statement; consult the official documentation using the links in the earlier footnote.

After inserting the values, a basic SELECT statement, which is the core querying command in SQL, checks that the data is actually in the tables. Run the commands in the following SQL code block one at a time to see each command's results:

```
SELECT * FROM products;
SELECT * FROM suppliers;
```

This is the simplest form of a SELECT statement, the asterisk ("*") instructs SQL to retrieve all columns. Later examples will illustrate ways to retrieve only some columns, and many other variants on the SELECT statement.

The two tables allow capturing information about products and information about suppliers, but they do not allow capturing which supplier supplies which product. In order to do this, the two tables need to be related by a foreign key relationship.

For the following example, assume that suppliers can supply many products, but a product may be supplied by only one supplier (or no supplier at all). This is called a *one-to-many relationship*. The following SQL code block alters the tables, retaining the existing data, and then updates the information in the new "supplier" column for the "products" table:

```
ALTER TABLE products ADD COLUMN supplier integer;
ALTER TABLE products ADD FOREIGN KEY (supplier) REFERENCES suppliers;

UPDATE products SET supplier = 1 WHERE pcode = 1;
UPDATE products SET supplier = 1 WHERE pcode = 2;
```

The first SQL statement above adds a new column to the existing products table in which to record the supplier of the product. The second line creates a foreign key reference from the supplier column in the products table to the primary key of the suppliers table; the primary is the "scode" column (see SQL code above). This ensures that only those suppliers can be recorded in the products table that actually exist in the suppliers table.

The third and fourth line update the data in the products table and set the value of the supplier column for different products. The two products have the same supplier which reflects the assumption that a supplier may supply multiple products. On the other hand, only one supplier can be recorded for each product, and this too reflects the above assumption. This expresses the *one-to-many relationship*. Moreover, the value of the supplier column in the products table may be NULL. In fact, after altering the table to add this column, all its values were NULL. A NULL value reflects the fact that a product has no supplier.

As an alternative to altering the existing products table, drop the products table to delete it and re-create it. Then insert some values. The following SQL code block uses the DROP TABLE command of SQL to delete the products table and all its contents.

```
DROP TABLE products;

CREATE TABLE products (
    pcode      integer,
    name       varchar(100),
    price      float4,
    supplier   integer,
    PRIMARY KEY (pcode),
    FOREIGN KEY (supplier) REFERENCES suppliers
);

INSERT INTO products VALUES(1, 'Hex Bolt', 1.99, 1);
INSERT INTO products VALUES(2, 'Round Bolt', 2.99, 1);
INSERT INTO products VALUES(3, 'Square Bolt', 3.99, NULL);
```

The above SQL code achieves the same as altering the table but in the process deletes all data in the products table. When possible, it is therefore preferable to use multiple ALTER TABLE statements instead of DROP and CREATE statements.

Note the following important points about the tables so far:

- There are products that have no supplier (the "square bolt")

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- There are suppliers that supply many products (supplier 1)
- There are suppliers that do not supply products (supplier 2)

In the products table as altered or re-created to this point, it is possible that a product has no supplier. However, in some applications it may be necessary to enforce that it is mandatory for products to have a supplier. This is done by adding a NOT NULL constraint, either by altering the table again, as in the following SQL code block, or by re-creating it with the appropriate constraint added.

```
ALTER TABLE products ALTER COLUMN supplier SET NOT NULL;
```

Adding constraints can only be done when the constraint is already satisfied. This means that in this example, none of the values of the supplier columns can be NULL when adding the constraint. If a new constraint is violated, the DBMS will show an error and the constraint will not be added.

When re-creating the table, the NOT NULL column constraint can be defined in the CREATE TABLE statement:

```
DROP TABLE IF EXISTS products;

CREATE TABLE products (
    pcode      integer,
    name       varchar(100),
    price      float4,
    supplier   integer NOT NULL,
    PRIMARY KEY (pcode),
    FOREIGN KEY (supplier) REFERENCES suppliers
);
```

So far, the assumption was that each product can have one supplier. However, in many settings, products have multiple suppliers, and suppliers supply multiple products, that is, there is a *many-to-many relationship* between the two. Expressing many-to-many relationships requires a third table that explicitly represents the relationship, here the "supplies" relationship between products and suppliers. The following SQL code first removes the existing tables, then re-creates tables to express a many-to-many relationship instead.

```

DROP TABLE IF EXISTS products;
DROP TABLE IF EXISTS suppliers;

CREATE TABLE products (
    pcode integer,
    name varchar(100),
    PRIMARY KEY (pcode) );

CREATE TABLE suppliers (
    scode integer,
    name varchar(100),
    city varchar(100),
    PRIMARY KEY (scode) );

CREATE TABLE supplies (
    scode integer NOT NULL,
    pcode integer NOT NULL,
    price float4 NOT NULL,
    PRIMARY KEY (scode, pcode),
    FOREIGN KEY (scode) REFERENCES suppliers,
    FOREIGN KEY (pcode) REFERENCES products );

```

Note that the tables must be dropped in the right order: "products" first, then "suppliers" because the products depend on the suppliers due to the foreign key constraint⁷. The IF EXISTS part is a safeguard to prevent an error if the table does not exist when attempting to drop it.

The primary key of the supplies table is a *compound key*, that is, it consists of a combination of columns. The supplies table is related by two FOREIGN KEY constraints both to the products and suppliers table so that only products and suppliers that already exist can be recorded here (and thereby related to each other). The price column is no longer in the products table, but has been moved to the supplies table, because each supplier may supply a product at a different price. The following example data shows this:

```

INSERT INTO products VALUES (1, 'Hex Bolt');
INSERT INTO products VALUES (2, 'Round Bolt');

INSERT INTO suppliers VALUES (1, 'Bolts Inc', 'HVGB');
INSERT INTO suppliers VALUES (2, 'Hardware Co', 'Cartwright');

INSERT INTO supplies VALUES(1, 1, 1.99);
INSERT INTO supplies VALUES(1, 2, 2.49);
INSERT INTO supplies VALUES(2, 1, 2.99);
INSERT INTO supplies VALUES(2, 2, 1.79);

```

To clean up after these exercises, drop all tables if they are no longer required:

⁷Use the CASCADE keyword to drop dependent tables automatically but use with care.

```
DROP TABLE supplies;
DROP TABLE products;
DROP TABLE suppliers;
```

Summary In summary, a *one-to-many relationships* requires a foreign key from the "many" table that references the "one" table and its primary key. In the first example above, a supplier supplies many products but a product has one supplier (or none, depending on whether a NOT NULL constraint has been specified). In contrast, a *many-to-many relationship* requires a table that explicitly represents the relationship. Foreign keys from this table reference the participating, original, "main" tables and their primary keys. In the second example above, a supplier supplies many products and a product can be supplied by many suppliers.

In fact, this type of relationship can be extended in a straightforward way to three or more tables. For example, a supplier supplies many products from many warehouses, a product may be supplied by many suppliers from many warehouses, and a warehouse may contain many products from many suppliers.

Hands-On Exercise

1. Consider the following information:
 - A book has an ISBN number and a title.
 - An author has a name and an address.
 - An author can write many books, and a book can be written by multiple authors. A book is written in a certain year.
2. Write the CREATE TABLE statements with the necessary FOREIGN KEY statements, and execute them on PostgreSQL
 - Use appropriate datatypes for the columns
 - Create an appropriate PRIMARY KEY for all tables
3. Use INSERT statements to create some example data.
4. Use SELECT statements to ensure your data exists.

3.5 SQL Queries

The previous section has presented the basics of the relational database model, focusing on how tables are related by foreign key relationships. Tables and their relationships are often graphically shown in a *relational diagram*. Such diagrams are often called "*ER Diagrams*"⁸. A graphical representation of the database structure is useful for understanding the data and for writing queries to extract data from the table or tables of the database.

⁸Technically, the two are not quite the same.

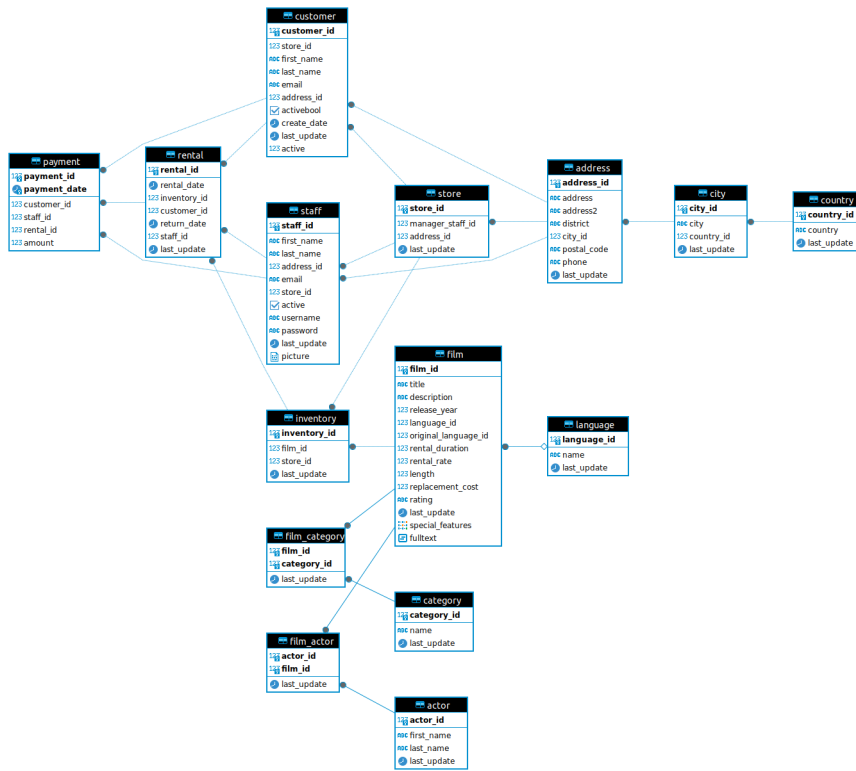


Figure 3.4: Relational diagram of the Pagila demo database

In many software tools, including in pgAdmin, the database developer can use relational diagrams to create tables, instead of writing CREATE TABLE statements. In the reverse, relational diagrams can also be automatically created from an existing database and its tables.

Hands-On Exercise

In the pgAdmin Object Explorer, right-click on the "busi4720" database, then select "ERD for Database" to create a relational diagram.

In the DBeaver Database Navigator, select the "busi4720" database, then right-click on its "public" schema and select "View Diagram" to create the diagram.

The resulting diagram will look similar to the one in Figure 3.4. (If you did not clean up the tables you created in the above SQL exercises, these will be present in the diagram as well).

The Pagila Database

The diagram shows the structure of the *Pagila* database⁹, a demonstration database originally developed for teaching and development of the MySQL RDBMS under the name *Sakila*¹⁰. *Pagila* is designed as a sample database to illustrate database concepts and is based on a fictional DVD rental store. It consists of multiple tables for film and actor information, customer data, store inventory, and rental transactions. Here is an overview of the key tables in the *Pagila* database:

- *actor*: Stores details about actors, including their first and last names.
- *film*: Contains information about movies, such as title, release year, language, rental duration, rental rate, length, replacement cost, rating, and special features.
- *film_actor*: A junction table that establishes a many-to-many relationship between the films and actors. It links each film to its actors.
- *category*: Lists different genres or categories of films.
- *film_category*: Another junction table that connects films to their respective categories.
- *language*: Stores languages in which the films are available.
- *customer*: Contains customer information, including names, email addresses, addresses, and store ID where they are registered.
- *address*: Holds address details, including city, postal code, phone number, and other address components.
- *city*: Contains information about cities, linked to the addresses.
- *country*: Stores country information, which is linked to cities.
- *store*: Includes data about the DVD rental stores, such as the store's address and the staff.
- *inventory*: Contains information about the store's inventory, including which film copies are available at which store.
- *rental*: Records details about rental transactions, including rental and return dates, inventory, and customer information.
- *payment*: Tracks payments made by customers for rentals, including amount and payment date.
- *staff*: Contains information about the store staff, including their names, email addresses, and the store they work in.

⁹<https://github.com/devrimgunduz/pagila>,
<https://github.com/devrimgunduz/pagila/blob/master/LICENSE.txt>

¹⁰<https://dev.mysql.com/doc/sakila/en/>,
<https://dev.mysql.com/doc/sakila/en/sakila-license.html>

Each table is designed with primary keys for unique identification and foreign keys to establish relationships with other tables. This structure allows for complex queries across multiple tables, facilitating a wide range of analyses, from inventory management to customer behavior tracking. The Pagila database is a good example of a real-world database schema and offers a good data set for practicing SQL queries.

The SELECT Statement

The SELECT statement in SQL is used to query and retrieve data from one or more tables in a database. The basic structure of a SELECT statement allows specification of which columns of data you want to retrieve and from which tables. A SELECT statement has multiple clauses or parts that are used to specify different characteristics of the information to retrieve:

- **SELECT:** Which columns to query (use the asterisk “*” to select all).
- **FROM:** Which tables to query from.
- **JOIN:** How to combine data from multiple tables based on related columns.
- **WHERE:** Conditions on field values used to filter the retrieved records.
- **GROUP BY:** Groups within which to aggregate data using an aggregate function such as `sum()`, `count()`, or `max()`.
- **HAVING:** Conditions on group aggregate values. Similar to WHERE but for aggregates within each group.
- **ORDER BY:** How to sort the resulting records in either ascending or descending order.
- **LIMIT:** Limit on how many results to return.

The following examples show queries for the Pagila database to illustrate different features of the SELECT statement. Instead of describing every option in detail, to understand SQL it is useful to execute the queries and learn by modifying the queries and observing changes in the results.

Example: Find all actors and the films they appeared in, ordered by film category and year, for those films that are rated PG

```

SELECT concat(left(actor.first_name, 1), '.', 
    actor.last_name) AS Actor,
    category.name AS Category,
    film.title,
    film.release_year
FROM film_actor
INNER JOIN actor USING (actor_id)
INNER JOIN film USING (film_id)
INNER JOIN film_category USING (film_id)
INNER JOIN category USING (category_id)
WHERE film.rating = 'PG'
ORDER BY actor.last_name,
    actor.first_name,
    category.name ASC,
    film.release_year DESC,
    film.title ASC;

```

Running this query will retrieve 1143 records from the Pagila database.

The SELECT clause specifies only a few columns to retrieve. Note that column names are prefixed by the table name, as in "actor.first_name" to make them unambiguous when multiple tables contain columns with the same name. Some columns are also given *aliases* using the keyword AS. This is useful to give the results more meaningful and shorter names. The first item to be selected is the result of a *function*: The concat() function concatenates text, and the left() function extracts the left part of some text. Refer to the PostgreSQL documentation linked to in the footnote above for a complete reference to available functions.

The FROM clause specifies a single table "film_actor" to which other tables are joined using the INNER JOIN keyword. The JOIN clause specifies the common join column with the USING keyword. The join columns typically correspond to the columns related by a foreign key relationship. In the Pagila database, foreign keys are always single columns, as are primary keys. However, one can imagine that combinations of two or more columns serve as primary keys and as foreign keys. Then, multiple columns are specified in the USING clause.

An INNER JOIN is a type of join that matches records from two tables if they both have the same value in their join columns, and only if they both have non-null values in their join columns. In contrast, a LEFT OUTER JOIN would also include records from the left table that have a NULL value in their join columns, a RIGHT OUTER JOIN would also include records from the right table that have a NULL value in their join columns, and a FULL OUTER JOIN is the combination of a LEFT OUTER JOIN and a RIGHT OUTER JOIN.

Compare the FROM and JOIN clauses to the relational diagram in Figure 3.4 and notice how it allows you to "navigate" from one table to another table along the foreign key relationships that link each table.

The WHERE clause in the above example selects those films whose rating is equal to

the text "PG". Multiple logical conditions can be combined with the AND, OR, NOT keywords and parentheses.

The ORDER BY clause specifies the ordering of the results. In this case, ordering is first done by actor last name. When actors have the same last name, ordering is done by first name. Within the same last and first names, ordering is done by category name, in ascending order, as indicated by the ASC keyword (The default ordering is always ascending, but it is sometimes useful to explicitly indicate this). Next, results are sorted by film release year in descending order, then again by film title in ascending order.

The JOIN ... USING clause assumes that the columns have the same name in both tables. When this is not the case, this short form is not available and joins must be specified manually. Recall that the join ensures that the join columns in both tables have the same value, which can also be specified in a WHERE clause as a set of conditions. The following query is equivalent to the previous one, but without the JOIN clauses:

```
SELECT concat(left(actor.first_name, 1), ' ',  
            actor.last_name) AS Actor,  
      category.name AS Category,  
      film.title,  
      film.release_year  
  FROM film_actor, film, actor, film_category, category  
 WHERE actor.actor_id = film_actor.actor_id AND  
       film.film_id = film_actor.film_id AND  
       film_category.film_id = film.film_id AND  
       category.category_id = film_category.category_id AND  
       film.rating = 'PG'  
 ORDER BY actor.last_name,  
          actor.first_name,  
          category.name ASC,  
          film.release_year DESC,  
          film.title ASC;
```

When writing the query without JOIN keywords, the required tables must all be included in the FROM clause; it does not matter in which order they are listed there.

Example: Find the most popular actors in the rentals in each city

```

SELECT city.city,
       concat(actor.first_name, ' ', actor.last_name) AS actor_name,
       count(rental.rental_id) AS Number_Rentals
  FROM rental
 INNER JOIN inventory USING (inventory_id)
 INNER JOIN store USING (store_id)
 INNER JOIN address USING (address_id)
 INNER JOIN city USING (city_id)
 INNER JOIN film USING (film_id)
 INNER JOIN film_actor USING (film_id)
 INNER JOIN actor USING (actor_id)
 GROUP BY city.city, actor.actor_id
 HAVING count(rental.rental_id) >= 300
 ORDER BY city ASC,
          Number_Rentals DESC,
          actor_name ASC;

```

Running this query will retrieve 22 records from the Pagila database.

This query uses the `GROUP BY` clause to group data. First, data is grouped by the city name, then, within each city, data is grouped by actor identification. Grouping allows, and in fact requires, the use of *aggregate functions*. This query uses the `count()` function to count the number of rentals in each group, that is, for each combination of city and actor. This query also includes a `HAVING` clause to return only those groups for which the count of rentals is greater than or equal to 300.

Example: Find the customers who spent the most, with their phone numbers and cities, the cities their store is in, and the number of rentals with the highest total rental payments for each film category, grouped by city of the rental store.

```

SELECT category.name AS category_name,
       store_city.city AS store_city,
       customer.customer_id,
       concat(customer.first_name, ' ',
              customer.last_name) AS customer_name,
       cust_city.city AS customer_city,
       cust_address.phone AS customer_phone,
       count(rental.rental_id) AS num_rentals,
       sum(amount) AS total_amount
  FROM city AS cust_city, city AS store_city,
       address AS cust_address, address AS store_address,
       store, rental
 INNER JOIN payment USING (customer_id)
 INNER JOIN customer USING (customer_id)
 INNER JOIN inventory USING (inventory_id)
 INNER JOIN film USING (film_id)
 INNER JOIN film_category USING (film_id)
 INNER JOIN category USING (category_id)
 WHERE store.store_id = inventory.store_id
   AND store_address.address_id = store.address_id
   AND store_city.city_id = store_address.city_id

```

```

        AND cust_address.address_id = customer.address_id
        AND cust_city.city_id = cust_address.city_id
GROUP BY category.name, customer.customer_id,
         cust_address.address_id, cust_city.city, store_city.city
HAVING sum(amount) IN (
    SELECT sum(amount) AS maxamount
        FROM store, address, city AS inner_city, rental
        INNER JOIN payment USING (customer_id)
        INNER JOIN customer USING (customer_id)
        INNER JOIN inventory USING (inventory_id)
        INNER JOIN film USING (film_id)
        INNER JOIN film_category USING (film_id)
        INNER JOIN category AS inner_category USING (category_id)
    WHERE inner_category.name = category.name AND
          inner_city.city = store_city.city AND
          store.store_id = inventory.store_id AND
          address.address_id = store.address_id AND
          inner_city.city_id = address.city_id
GROUP BY inner_category.name, inner_city.city,
         customer.customer_id
ORDER BY inner_category.name ASC, inner_city.city,
         maxamount DESC
LIMIT 1 )
ORDER BY category.name ASC, store_city ASC;

```

This query will return 33 records from the Pagila database.

This complex query is actually two queries as it includes a *subquery* within the HAVING clause! Starting in the FROM clause, notice that some tables are included twice in this query, under different *aliases* or names, using the AS keyword. This is because the query retrieves the city that the customers live in, as well as the city that the store is located in. As cities are linked to addresses, the address table is also included twice.

Some joins are done using the JOIN keyword on common columns, while joining the address and city tables is done in the WHERE clause because the column names "city_id" and "address_id" are ambiguous as the tables are included multiple times.

The GROUP BY keyword groups the results by category, customer, address, customer city, and store city. This is necessary to be able to the select the customer city and address: Only columns that are grouped by can be retrieved or selected and aggregated, for example by using the sum() function used in this query.

As the query seeks to retrieve the maximum amount spent, the HAVING clause is used to select just this maximum by ensuring that the sum of payment amounts is equal to the result of the subquery. This subquery is very similar to the "outer" query, but returns only the sum of payment amounts, ordered by this amount in descending order and limited to the first result. That is, the subquery returns the maximum sum of payment amounts. The subquery is linked to the outer query by two conditions in its WHERE clause: inner_category.name = category.name and inner_city.city == store_city.city. These two conditions ensure that

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the maximum sum of payments computed in the subquery is done for the same grouping that the outer query is considering.

Because of the subquery, running this query is expensive in terms of computing time.

Example: Get the total rental revenue and number of rentals for each store by month

```
SELECT city.city,
       extract(year from payment_date) AS year,
       extract(month from payment_date) AS month,
       sum(amount) AS dollars,
       count(rental.rental_id) AS rentals
  FROM payment, rental, inventory, store, address, city
 WHERE payment.rental_id = rental.rental_id AND
       rental.inventory_id = inventory.inventory_id AND
       inventory.store_id = store.store_id AND
       store.address_id = address.address_id AND
       address.city_id = city.city_id
 GROUP BY city.city,
          extract(year from payment_date),
          extract(month from payment_date)
 ORDER BY city.city,
          extract(year from payment_date),
          extract(month from payment_date);
```

Running this query will return 14 results.

This query shows the use of a date function in PostgreSQL. The `extract()` function can extract part of a date. In the example, it is used to extract the year and the month. Both are also used in the `GROUP BY` and in the `ORDER BY` clause.

Example: Get the top 5 and the bottom 5 grossing customers by year

```
( SELECT concat(customer.first_name, ' ',
                customer.last_name) AS customer_name,
      extract(year from payment_date) AS year,
      sum(amount) AS dollars,
      'Top-5' AS note
  FROM payment, customer
 WHERE payment.customer_id = customer.customer_id
 GROUP BY extract(year from payment_date),
          customer.customer_id
 ORDER BY dollars DESC
 LIMIT 5
) UNION (
SELECT concat(customer.first_name, ' ',
                customer.last_name) AS customer_name,
      extract(year from payment_date) AS year,
      sum(amount) AS dollars,
      'Bottom-5' AS note
  FROM payment, customer
 WHERE payment.customer_id = customer.customer_id
 GROUP BY extract(year from payment_date),
          customer.customer_id
 ORDER BY dollars ASC
LIMIT 5 ) ORDER BY dollars DESC;
```

This query combines the results of two simple queries with the UNION keyword. Both queries must return the same columns in order to be combined in this way. Because the results are mathematically sets, they are not intrinsically ordered; this is why the set that results from the UNION operation is ordered again.

Set operations can be used to combine results from multiple queries. These are specified by the UNION, INTERSECT, and EXCEPT keywords and do exactly as their name indicates: They return the union, the intersection, or the complement of two result sets. The inputs to each operation sets must have the same set of columns.

PostgreSQL can easily import and export data for further analysis. The pgAdmin application can export query results to CSV files (there is a button in the query toolbar). Alternatively, one can use the *COPY* command as in the following example.

```
COPY (SELECT * FROM customer)
 TO '/tmp/filename.csv'
 WITH (FORMAT CSV, HEADER);
```

Similarly, PostgreSQL can easily import data from CSV files using the copy command:

```
COPY customer
 FROM '/tmp/filename.csv'
 WITH (FORMAT CSV, HEADER);
```

Additionally, PostgreSQL can import and export JSON files; see the documentation for details.

Hands-On Exercise

1. Find the names and the rental numbers of the top 5 customers who rented the most films
 - **Tip:** Join tables "rental", "customer", use the "count()" function
2. Calculate the rental revenue per customer. Who are the top 5? Bottom 5?
 - **Tip:** Join tables "rental", "customer", "payment", use the "sum()" function
3. Calculate the average rental revenue per customer for each store
 - **Tip:** Join tables "rental", "customer", "payment", "inventory", use the "avg()" function
4. Calculate the rental counts for each country of customer. Are there countries with no rentals?
 - **Tip:** Join tables "rental", "customer", "address", "city", "country", use the "count()" function
5. Find all films with a single actor
 - **Tip:** Join tables "film", "film_actor", use the "count()" function in a HAVING clause
6. Create tables to represent a part-of hierarchy. For example, a product may be a part of another product, and products may have multiple parts.
 - **Tip:** You need only one table

3.6 Review Questions

1. What is a relational database, and who developed the relational model?
2. Explain the role of primary keys and foreign keys in relational databases.
3. What is SQL and what are its main purposes?
4. List and describe at least four data types commonly used in SQL.
5. Explain the difference between the `varchar` and `text` data types in PostgreSQL.
6. What are the ACID properties in relational databases and what is their purpose?
7. Define and give an example of each of the following constraints:
 - A. NOT NULL
 - B. UNIQUE
 - C. PRIMARY KEY
 - D. FOREIGN KEY
 - E. CHECK
8. How do relational databases handle relationships between tables? Give examples.
9. What are some of the challenges relational databases faced with the advent of Big Data?

10. What is PostgreSQL and what type of system is it?
11. What is the purpose of the “psql” and “pgAdmin” tools in the context of PostgreSQL?
12. When connecting to a DBMS running on your own computer, what hostname should you use?
13. Define a “schema” in the context of a PostgreSQL database. What is the default schema in PostgreSQL?

3.7 Additional SQL Exercises

Database Schema:

- **Table:** Employees
- **Columns:** EmployeeID, FirstName, LastName, Role, Department

Task: Write a SQL query to select the first and last names of all employees in the ‘Sales’ department.

Database Schema:

- **Table:** Products
- **Columns:** ProductID, ProductName, Price, Category, StockQuantity

Task: Write a SQL query to select the ProductName and Price for all products in the ‘Electronics’ category.

Database Schema:

- **Table:** Books
- **Columns:** BookID, Title, Author, PublishYear, Price

Task: Write a SQL query to select all columns from the Books table and sort the results by PublishYear in descending order.

Database Schema:

- **Table:** Orders
- **Columns:** OrderID, CustomerName, OrderDate, TotalAmount

Task: Write a SQL query to select the OrderID and TotalAmount for orders where the TotalAmount is greater than 100. Sort the results by TotalAmount in ascending order.

Database Schema:

- **Table:** Students

- **Columns:** StudentID, Name, Major

Task: Write a SQL query to select all distinct majors from the Students table.

Database Schema:

- **Table:** Customers
- **Columns:** CustomerID, FirstName, LastName, Email
- **Table:** Orders
- **Columns:** OrderID, CustomerID, OrderDate, TotalAmount

Task: Write a SQL query to select all orders with the corresponding customer's first and last name. Join the Customers and Orders tables on CustomerID.

Database Schema:

- **Table:** Authors
- **Columns:** AuthorID, Name
- **Table:** Books
- **Columns:** BookID, Title, AuthorID
- **Table:** Publishers
- **Columns:** PublisherID, Name
- **Table:** BookPublishers
- **Columns:** BookID, PublisherID

Task: Write a SQL query to select the title of the book, the name of the author, and the name of the publisher. This will require joining the Books, Authors, and BookPublishers tables, and then joining the resulting table with Publishers.

Database Schema:

- **Table:** Employees
- **Columns:** EmployeeID, FirstName, LastName, Department
- **Table:** Sales
- **Columns:** SaleID, EmployeeID, SaleAmount, SaleDate

Task: Write a SQL query to select each employee's first name, last name, and total sales amount. This requires a join between Employees and Sales tables and the use of the SUM aggregate function on SaleAmount.

Database Schema:

- **Table:** Products
- **Columns:** ProductID, ProductName, Price
- **Table:** Orders
- **Columns:** OrderID, ProductID, Quantity

Task: Write a SQL query to select all products, along with the quantity ordered for each product. Use a LEFT JOIN to ensure that all products are listed, even if they have not been ordered.

Database Schema:

- **Table:** Students
- **Columns:** StudentID, Name, Major
- **Table:** Enrollments
- **Columns:** CourseID, StudentID, Grade

Task: Write a SQL query to select the names of students and their grades who are enrolled in a specific course (e.g., 'Biology 101'). This requires a join between the Students and Enrollments tables and a WHERE clause to filter by the CourseID.

Database Schema:

- **Table:** Employees
- **Columns:** EmployeeID, FirstName, LastName, Salary, DepartmentID
- **Table:** Departments
- **Columns:** DepartmentID, DepartmentName

Task: Write a SQL query to select the first name and last name of employees who earn more than the average salary in their respective departments. This will require a subquery in the WHERE clause to calculate the average salary per department.

Database Schema:

- **Table:** Movies
- **Columns:** MovieID, Title, ReleaseYear, Genre
- **Table:** Ratings
- **Columns:** RatingID, MovieID, Reviewer, Stars

Task: Write a SQL query to select the title of movies that have an average rating higher than the overall average rating of all movies. This will require a complex subquery to first calculate the average rating for each movie, and another subquery to calculate the

overall average rating.

Chapter 4

Managing Graph Data with Graph Databases

4.1 Introduction

Graph databases are one type of *NoSQL* databases, an acronym for "Not Only SQL". NoSQL databases emerged as a response to the limitations of traditional relational database systems and the evolving needs of modern applications. The concept and the term "NoSQL" gained prominence in the late 2000s, but its roots can be traced back to earlier innovations in database technology.

The rise of the internet and web applications in the 1990s and 2000s led to unprecedented amounts of data and new types of data that did not fit neatly into the rows and columns of relational databases. Companies like Google and Amazon faced challenges in scaling their databases to meet the demands of huge amounts of web traffic and large, unstructured data sets. This led to the development of new database systems like Google's Bigtable and Amazon's Dynamo, which laid the groundwork for NoSQL databases.

NoSQL databases were designed to overcome the scalability, performance, and flexibility limitations of traditional relational databases. Unlike relational databases that use a fixed table structure, NoSQL databases utilize a variety of data models, including key-value, document, and graph formats. This diversity allows them to handle a wide array of data types and structures efficiently.

Key benefits of NoSQL databases include their ability to scale horizontally across many servers, offering significant advantages in handling large-scale, high-volume applications and big data. They also deliver high performance, particularly with large volumes of data and concurrent read/write operations, due in part to their typical emphasis on eventual data consistency over strict ACID compliance (atomicity, consistency, isolation, and durability of database transactions).

The schema-less nature of NoSQL databases provides more agility in application development. Developers can iterate quickly without needing to restructure databases every time the application evolves. This flexibility is especially valuable in agile software development environments and for applications dealing with diverse, unstructured, or rapidly evolving data sets.

Moreover, many NoSQL databases natively support modern data formats like JSON, aligning well with current web and mobile applications. This can simplify the development process, as the same data format can be used throughout the application stack.

However, the strengths of NoSQL databases also bring some drawbacks. The lack of a fixed schema means that data integrity cannot be ensured using typing of columns, primary keys on unique identifiers, or referential integrity with foreign keys. NoSQL databases also do not typically make correctness guarantees for concurrent transactions that come from the ACID properties of relational databases. Instead, they guarantee that eventually the data will be consistent, but applications and users may occasionally see inconsistent data. Generally NoSQL databases are less suitable for high-volume concurrent update transaction processing; these application types are better supported by relational databases. Instead, NoSQL databases are better suited for applications that may require complex queries but relatively infrequent updates, few concurrent update transactions, and updates of single data elements at a time.

Graph databases are designed to store and query relationships in data. They represent data as nodes, akin to entities in a relational database, and relationships between these nodes. This structure is particularly suited for handling complex, interconnected data and is highly efficient in scenarios where relationships are as important as the data itself. Graph databases gained significant traction driven by the increasing complexity of data and the limitations of relational databases in efficiently handling highly connected or networked data. The proliferation of social networks, recommendation systems, and other applications dealing with complex relationships between data entities spurred the development of graph databases.

Unlike relational databases that require computationally intensive join operations to establish connections between data in different tables, graph databases are designed to store relationships as first-class objects. This means that queries on interconnected data are faster and more efficient, as they exploit the direct connections between nodes. Additionally, graph databases are schema-less or have flexible schemas, allowing for more agility in adapting to changing data requirements.

4.2 Use Cases

Graph databases have become increasingly important in various industries due to their ability to efficiently model and query complex relationships and interconnected data that arise in those applications.

In *fraud detection*, graph databases are used to uncover patterns that are indicative of fraudulent activities. They can map complex transaction networks and identify un-

usual patterns, such as circular transactions or abnormally close relationships between entities, that might signal fraud. The ability to quickly traverse and analyze complex networks of data helps in real-time detection and prevention of fraud.

For *IT infrastructure monitoring*, graph databases offer a way to model complex networks of servers, devices, and applications. They can track the relationships and dependencies between various components of an IT system. This is invaluable for root cause analysis, where understanding the impact of an issue in one part of the system on the rest is crucial for quick resolution.

Graph databases power *recommender engines* by capturing and analyzing relationships between users, their preferences, and products. They can efficiently traverse these relationships to generate personalized recommendations based on a user's past behavior and the behavior of similar users.

In *social media*, graph databases are used to model the complex relationships between users, their friends, and their activities. They help in understanding social dynamics, optimizing content delivery, and enhancing user engagement by providing insights into how users are connected and how information flows through these networks.

For *supply chain management*, graph databases can model the entire supply chain network, including suppliers, production facilities, distribution centers, and retail outlets. This aids in optimizing routes, managing inventories, and identifying vulnerabilities in the supply chain, such as single points of failure.

In the *financial sector*, graph databases are utilized for risk assessment, compliance, customer service, and understanding client relationships. They help in mapping and analyzing complex networks of transactions and customer relationships, which is critical for identifying risks, ensuring compliance with regulations, and offering personalized financial services.

In *life sciences*, graph databases play a significant role in drug discovery, genomics, and protein analysis. They are used to model complex biological systems and relationships, such as gene interactions, protein pathways, and patient data, assisting in research and the development of personalized medicine.

In each of these domains, the key advantage of graph databases lies in their ability to naturally represent complex networks and relationships. This allows for more intuitive data modeling, faster querying, and the extraction of insights that would be difficult or impossible to obtain with traditional relational databases.

4.3 Graph Database Languages

In contrast to the standardized SQL language for relational databases, graph databases use various query languages designed to leverage their unique structure and efficiently handle complex queries that involve interconnected data. One of the more prominent ones is the Cypher language, introduced and primarily used in the Neo4j graph database system since 2011 but opened for use in other systems in 2015 as the openCypher

project. It is a declarative language¹, known for its expressive and readable syntax tailored for describing patterns in graphs. Cypher allows for easy querying of nodes, relationships, and paths and includes powerful features for filtering, pattern matching, and aggregating data.

Another notable query language is Gremlin, part of the Apache TinkerPop graph computing framework. Development began in 2009 and is ongoing. Gremlin is versatile and functional, allowing for imperative and declarative querying across different graph databases. It is known for its flexibility and ability to execute both simple and complex traversals, making it suitable for a wide range of applications.

SPARQL (a recursive acronym for "SPARQL Protocol and RDF Query Language") is a query language used primarily for querying RDF (Resource Description Framework) data, often found in semantic web applications. Its development is overseen and standardized by the W3C (World Wide Web Consortium), beginning in 2008 with a major update in 2013. It is particularly suited to querying and manipulating data stored in RDF format, and is widely used in applications that require linking diverse data sources, such as knowledge graphs.

Additionally, some graph databases support SQL-like query languages with extensions to handle graph-specific structures. These languages make it easier for users familiar with SQL to transition to graph databases. An example is GraphQL, developed by Facebook in 2015.

The lack of a standard query language in the graph database realm has led to fragmentation. This fragmentation can pose challenges for users and developers, such as a steeper learning curve and difficulty in transitioning between different graph database systems. In response, the forthcoming standardized GQL (Graph Query Language) is a new graph query language specifically designed for interacting with graph databases. The development of GQL is overseen by ISO/IEC JTC 1, the same joint technical committee responsible for the SQL standard. Its design is expected to draw on the strengths of existing languages, offering robust features for graph traversal, pattern matching, and manipulation of graph structures while maintaining readability and ease of use. A first version of the GQL standard was expected for 2023.

4.4 The Neo4J Graph Database Management System

Cypher is the query language for Neo4j, one of the most popular graph database systems. It was specifically designed for querying the graph data in Neo4j, making it easy to work with complex graph structures. Cypher's syntax is intuitive and expressive, focusing on the clarity of graph patterns and drawing inspiration from SQL and other declarative query languages. Its pattern matching approach was styled after the SPARQL language. Key characteristics and features of Cypher include:

- *Graph Pattern Matching:* Cypher provides the ability to expressively describe

¹A declarative query language allows the user to specify *what* data to retrieve. In contrast, an imperative/procedural query language requires the user to specify *how* to retrieve data.

Getting Started	https://neo4j.com/docs/getting-started/
Cypher Manual	https://neo4j.com/docs/cypher-manual
Graph Data Science	https://neo4j.com/docs/graph-data-science
APOC Library	https://neo4j.com/docs/apoc/current/
Use Cases	https://neo4j.com/use-cases/
Resources	https://neo4j.com/resources/

Table 4.1: Neo4J Documentation

graph patterns. It uses a syntax where nodes and relationships in the graph are depicted using parentheses (representing nodes) and arrows (representing relationships). This makes it visually intuitive to understand the queries and the graph patterns they represent.

- *Rich Filtering Capabilities:* Cypher includes robust filtering capabilities, enabling users to write queries that can filter nodes and relationships based on various criteria, including properties and patterns.
- *Aggregation and Sorting:* Like SQL, Cypher allows for aggregating data, performing calculations, and sorting results. It provides functions for counting, summing, averaging, and other common aggregations.
- *Pathfinding and Graph Algorithms:* Cypher can handle common graph queries such as shortest path, reachable nodes, and more.
- *Subqueries and Joins:* Cypher supports subqueries and various forms of joins, enabling complex queries that can span multiple parts of the graph.
- *Extensibility:* Cypher can be extended with user-defined procedures and functions, allowing for custom logic and advanced processing capabilities.

Similar to SQL queries, Cypher queries have multiple clauses, specifying a "query pipeline" for selecting, filtering, and sorting data. Unlike SQL, Cypher queries allow graph reading and graph updating in the same Cypher statement.

Neo4J offers a number of options for running the Neo4J database management, among them a limited developer version called "Neo4J Desktop" and a free, open-source community edition that is usually accessed through a web interface ("Neo4J Browser"). Table 4.1 provides links to useful documentation of the Neo4J database and the Cypher language.

The community edition is installed in the course virtual machine and enabled to run when the machine is started. You can access Neo4J Browser (Figure 4.1) at <http://localhost:7474> with the username "**neo4j**" and the password "**busi4720**".

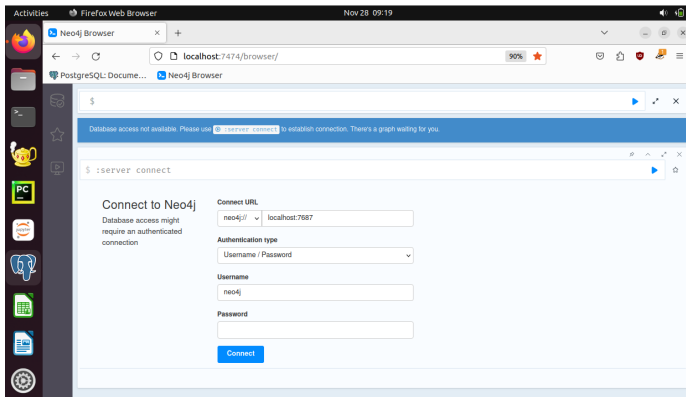


Figure 4.1: Neo4J Browser interface

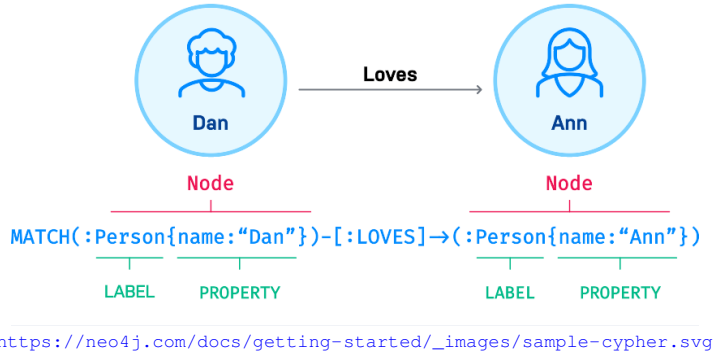


Figure 4.2: Sample Cypher syntax

4.5 Introduction to Cypher

Nodes Figure 4.2 shows how nodes and relationships are represented in the Cypher syntax. In Neo4J, nodes may be labelled with zero, one, or more labels. Labels are not types; a label does not specify anything about the information associated with a node, it merely serves to categorize or classify nodes. Nodes may have properties, specified as key–value pairs in JSON syntax. Graph nodes are written with normal round parentheses, with the set of their properties in curly brackets. Both the variable name for the node and the node labels are optional.

```
(variable : Label1:Label2:Label3 ... {k1:v1, k2:v2, k3:v3 ....})
```

Relationships Relationships are directed connections between two nodes. Unlike nodes, relationships only have a single label. But like nodes, they can have properties

specified as key–value pairs in JSON syntax. Relationships are written as "lines" between nodes, directed or undirected, with an optional variable name and relationship label in square brackets.

```
// Undirected, used in pattern
()-[variable : Label]-()
// Directed
()-[variable : Label]->()
// Directed
()<-[variable : Label]-()
// Unlabelled, no variable
() -[]-()
() -->()
() <--()
```

Only directed relationships can be created in a Neo4J graph, but undirected relationships can be used in a pattern to query a graph.

Directionality of relationships is important and matters for querying a graph. A directed relationship \rightarrow will match a directed pattern \rightarrow or an undirected pattern $-$ but not $<-$.

Path A path in Neo4J is a sequence of alternating nodes and relationships, beginning and ending with a node.

Patterns and Querying Graph patterns are used with the MATCH query keyword and describe either a node or a path that is to be searched for in the graph. When an instance of a pattern is found, any variable names in the pattern are bound to the corresponding nodes and relationship in the graph and the bound pattern is returned in the result set.

For example, consider the following simple pattern matching query. The MATCH clause specifies the pattern to match. The pattern here is a node, indicated by the use of round parentheses, and only a variable name n is specified, no labels or property values. This means that this pattern matches all nodes in the graph database and returns them in the variable names n .

```
MATCH (n)
```

The next pattern matching query adds a label Person to the node specification. This means the pattern matches only those nodes that are labelled as Person and returns them in the variable named p .

```
MATCH (p:Person)
```

Patterns can include property values to match. The following pattern matches all Person nodes that contain an attribute name with value 'Joe' and returns them in the variable p.

```
MATCH (p:Person {name: 'Joe'})
```

4.6 Defining Graphs in Cypher

Nodes or relationships in a graph can be created using the MERGE or CREATE statements in Cypher. As the name indicates, CREATE will create a node or relationship. In contrast, MERGE will check whether the node or relationship exists and only create it when it does not yet exist in the graph. Consider the example graph shown in Figure 4.3. The following Cypher codes creates this graph:

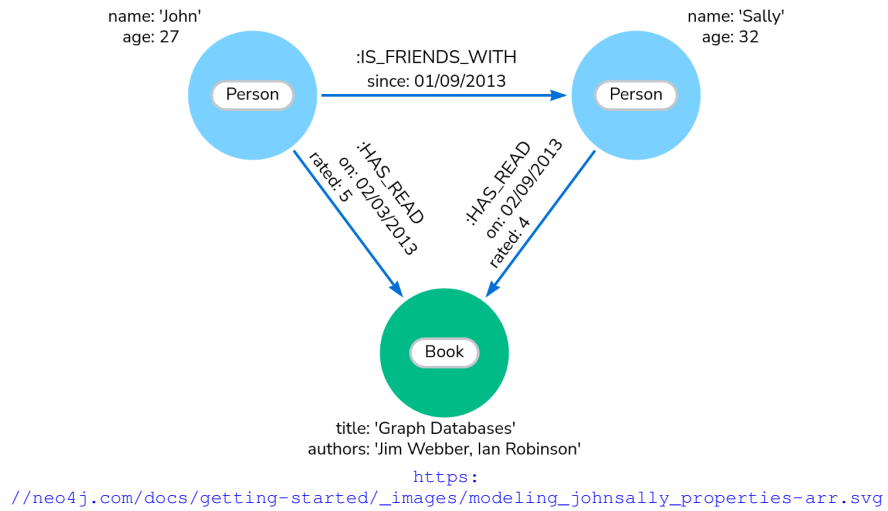


Figure 4.3: Example graph

```
// Create nodes
MERGE (j:Person {name: "John"})
  ON CREATE SET j.age = 27
MERGE (s:Person {name: "Sally"})
  ON CREATE SET s.age = 32
MERGE (b:Book {title: "Graph Databases"})
  ON CREATE SET b.authors = ["Jim Webber", "Ian Robinson"]

// Create relationships
MERGE (j)-[rel1:IS_FRIENDS_WITH]-(s)
  ON CREATE SET rel1.since = "01/09/2013"
MERGE (j)-[rel2:HAS_READ]-(b)
  ON CREATE SET rel2.on = "02/03/2013", rel2.rated = 5
MERGE (s)-[rel3:HAS_READ]-(b)
  ON CREATE SET rel3.on = "02/09/2013", rel3.rated = 4
```

Note that the six MERGE statements in the above code block are logically related, so that variable names, for example `j`, `b` and `s`, in one MERGE clause can be used to refer to a new node or relationship in a later MERGE clause. While using variable names in a MERGE clause is not mandatory, it is more efficient than having to later query the graph data for a particular node when creating subsequent relationships. The `ON CREATE SET` clause in the above statements sets one or more property values (separated by commas) of the newly created nodes and relationships. Note that some properties are lists, such as the `authors` property, indicated by the square brackets of the JSON notation.

The following MATCH queries can be used to view all nodes and relationships, irrespective of their labels. The first MATCH clause matches all nodes and returns them, the second MATCH clause matches all relationships between any two nodes and returns the relationships, the final query matches any two nodes that are connected by a relationship and returns the set of triples of first node, second node, and relationship.

```
// Query nodes
MATCH (n) RETURN n

// Query relationships
MATCH ()-[r]-() RETURN r

// Query both together
MATCH (n1)-[r]-(n2) RETURN n1, r, n2
```

The Neo4J Browser interface allows graph visualization and visual exploration of nodes, relationships, and their properties, as shown in Figure 4.4.

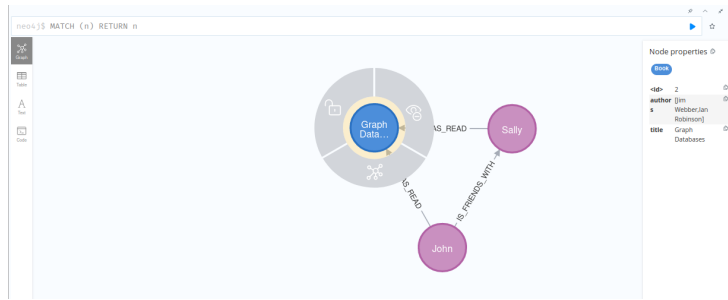


Figure 4.4: Graph Visualization and Exploration in Neo4J Browser

Hands-On Exercise

Consider the following description:

"You are completing the course BUSI 4720 in this semester with a final grade of 100. BUSI 4720 is part of the BCom program where it is offered in the 4th year. BUSI 4720 carries 3 credit hours of academic credit. It is a course on the topic of Business Analytics."

Define a graph in Cypher that represents this description:

1. Identify nodes, relationships, and properties of nodes and relationships
2. Use CREATE or MERGE statements to create nodes first, then relationships
3. Use MATCH to verify your graph is correct.

To remove nodes and relationships from a graph, use the MATCH query clause together with a DELETE clause. For example, to clean and remove the Person and Book nodes and relationships between Person and Book nodes created in the previous exercise, use the following Cypher statements:

```
MATCH (:Person|Book)-[r]-(:Person|Book) DELETE r;
MATCH (n:Person|Book) DELETE n;
```

To remove *all* relationships and nodes, irrespective of their label, omit node or relationship labels, as in the following Cypher code block. Use with care as this deletes all data in the graph database.

```
MATCH ()-[r]-() DELETE r;
MATCH (n) DELETE n;
```

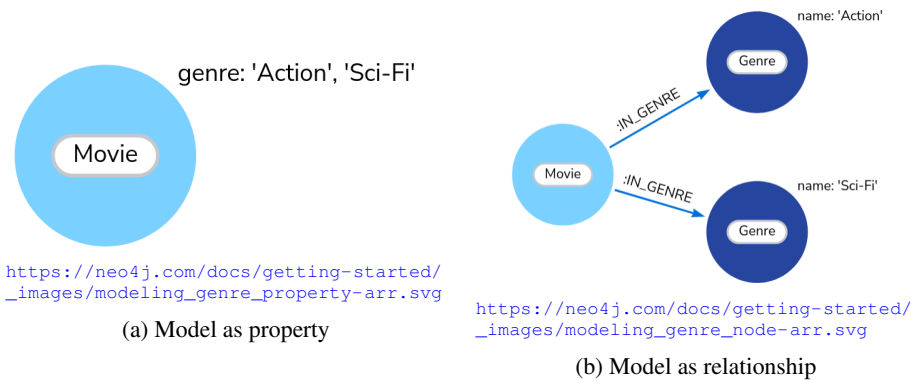


Figure 4.5: Equivalent graph models of movie genres

4.7 Graph Data Modeling

When defining a graph, one frequent question is whether to model something as a property of a node or as a relationship to a node. While there is no generally right or wrong answer to this question, the choice of data model depends on the queries to be run against the data, that is, the type of questions that will be asked.

Nodes versus Relationships

Consider the two graph models in Figures 4.5a and 4.5b. Both depict the same fact, that there exists a movie with title "The Matrix" in two genres, "Action", and "Sci-Fi". Figure 4.5a models the genres as a property of list type, that contains multiple entries in the list. In contrast, Figure 4.5b models the genres as nodes and the fact that the movie is in a genre as a relationship between the movie node and a genre node.

The graph model in Figure 4.5a is particularly useful to find the genres for a particular movie, that is, it is useful for queries that focus on the nodes and their properties. However, this model makes it difficult, cumbersome, and inefficient to find movies that share genres. The system has to consider all pairs of movies, and then for each pair of movies iterate through each of their property lists. The following two queries exemplify this. The first query simply filters the Movie nodes for a particular title and returns the genre attribute of the Movie node.

The second query first identifies all pairs of Movie nodes in the MATCH clause, then uses the WHERE clause to filter those pairs that share entries in their genre attribute. Recall that the genre attributes are lists, so `x IN m1.genre WHERE x IN m2.genre` checks every element of the second list for every element of the first list.

```
// find the genres for a particular movie
MATCH (m:Movie {title:"The Matrix"})
RETURN m.genre;

// find which movies share genres
MATCH (m1:Movie), (m2:Movie)
WHERE any(x IN m1.genre
          WHERE x IN m2.genre)
      AND m1 <> m2
RETURN m1, m2;
```

The graph model in Figure 4.5b on the other hand requires a more complex query to find the genres of the movie. However, while more complex, it is no less efficient than the corresponding query for the other model above. On the other hand, the query to find movies that share genres becomes easier, more intuitive to write, and more computationally efficient, as the following Cypher queries show.

The first query uses MATCH to first select movies and filter on the movie name, then for that movie `m` it traverses the `IN_GENRE` relationship to identify all related Genre nodes `g` in order to return their names. The second query is more intuitive than the corresponding query for the other model. It finds two Movie nodes `m1` and `m2` that both have an `IN_GENRE` relationship that points to the same Genre node `g`.

```
// find the genres for a particular movie
MATCH (m:Movie {title:"The Matrix"}),
      (m)-[:IN_GENRE]->(g:Genre)
RETURN g.name;

// find which movies share genres
MATCH (m1:Movie)-[:IN_GENRE]->(g:Genre),
      (m2:Movie)-[:IN_GENRE]->(g)
RETURN m1, m2, g
```

In summary, neither way of modeling the facts is better or worse, but the two options are more suitable to different types of queries and data to be retrieved.

Labels versus Attributes

Consider the two graph models in Figures 4.6a and 4.6b. The two models demonstrate the flexibility of modeling connected data and using labels to simplify queries and make them more efficient.

Figure 4.6a shows Airport nodes connected by `:FLYING_TO` relationships that indicate that a flight exists from one to the other airport. Information about flights is modelled as properties of the relationship. However, noting that multiple flights may be offered each day, it is clear that a flight node is required to represent each of those flights, with a node property that represents the date of the flight. However, when

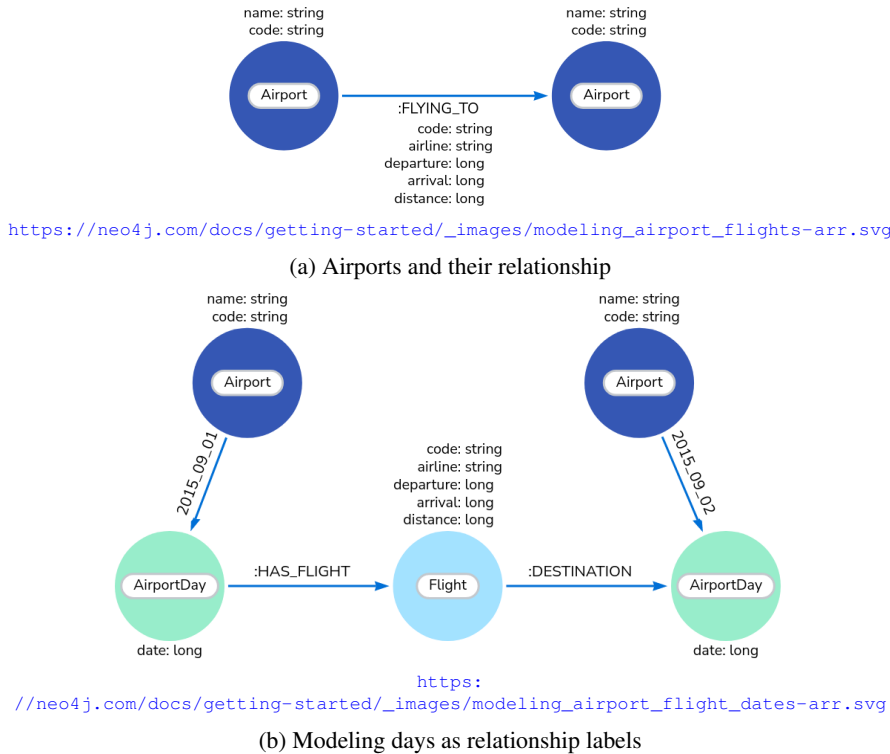


Figure 4.6: Graph models of airports and flights

querying such a model for flights on a particular date, the system must examine all flight nodes, and then filter those with the appropriate property value for the data.

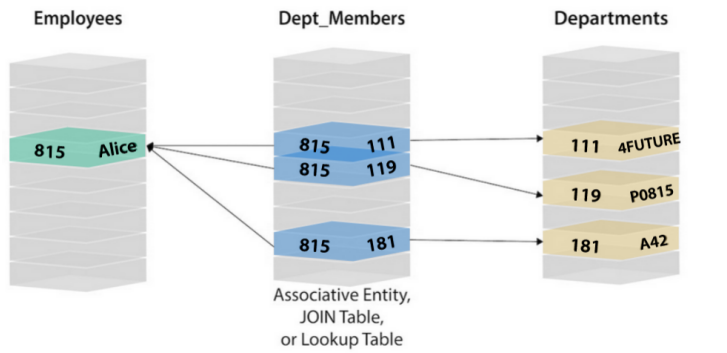
A more efficient model is that shown in Figure 4.6b. Here, the date of the flight is modelled as a label for the relationship between `Airport` and `AirportDay` nodes, which allows the system to easily select only those flights that occur on a certain date without having to examine all flight nodes.

This example shows again that the queries to be run against the data have a strong impact on how best to model your data, here affecting the decision whether to model data as an attribute or a label.

Relational Model and Graph Model

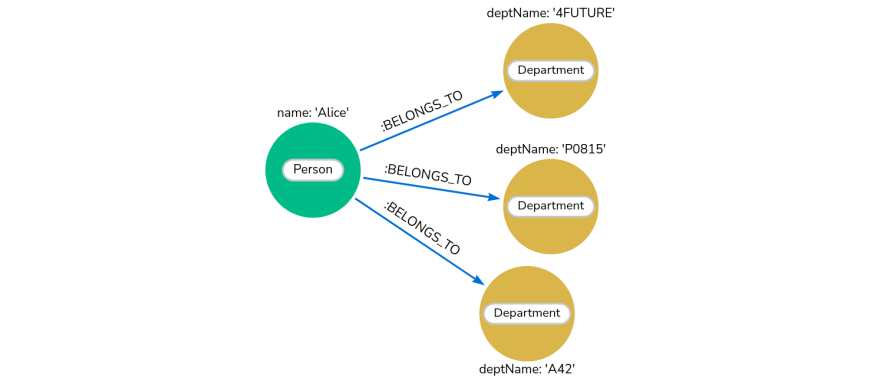
The relational data model consists of tables, their columns, and foreign key relationships that link tables (Figure 4.7a). It is straightforward to translate such a model to a graph model using the following translation heuristics:

- Table names become node labels



https://neo4j.com/docs/getting-started/_images/relational_model.svg

(a) A relational model



https://neo4j.com/docs/getting-started/_images/relational_graph_model-arr.svg

(b) An equivalent graph model

Figure 4.7: Transforming relational data to graph data

- Rows of data become nodes
- Columns become node properties
- Foreign keys become relationships between nodes
- Join tables become relationships between nodes; their properties become relationship properties
- Null values do not become properties, they are omitted entirely

Applying these heuristics to the example in Figure 4.7a leads to the graph model in Figure 4.7b. The table names "Employee" and "Department" have become node labels for two different categories of nodes. Each row in the employee table (for example, employee 815 with name Alice) is represented as a node with label **Person**, and each department (for example, department 111 with name 4Future) is a node with label **Department**. The column names, the "name" column in the Employees table and

the "deptName" column in the Departments table, have become properties of the corresponding nodes. The "Dept_Members" table joins employees and departments and has been transformed into the relationship with label BELONGS_TO between Person and Department nodes. The "Dept_Members" table had no columns other than those participating in the foreign key relations, but if it had, those columns would be attributes on the BELONGS_TO relationship.

Applying these heuristics should only be considered as an initial translation. As seen above, some or all properties may well be represented as nodes in their own right (Figure 4.5) or be modelled as relationship labels (Figure 4.6), depending on the type of queries expected to be run against the graph data.

Pagila Database Example As an example, each table of the Pagila relational database from the previous chapter was exported from PostgreSQL to a CSV file. These CSV files can be imported into Neo4J with the following set of Cypher expressions. Note that not all data is imported in this example, and a more compact representation of the statements is possible.

The Pagila database is already imported into the Neo4J Community Edition in the course virtual machine.

```

load csv with headers from 'file:///actor.csv' as row
merge (actor:Actor {actorID: row.actor_id})
on create set actor.firstName = row.first_name
on create set actor.lastName = row.last_name;

load csv with headers from 'file:///address.csv' as row
merge (address:Address {addressID: row.address_id})
on create set address.address = row.address
on create set address.district = row.district
on create set address.postalCode = row.postal_code
on create set address.phone = row.phone;

load csv with headers from 'file:///category.csv' as row
merge (category:Category {categoryID: row.category_id})
on create set category.name = row.name;

load csv with headers from 'file:///city.csv' as row
merge (city:City {cityID: row.city_id})
on create set city.city = row.city;

load csv with headers from 'file:///country.csv' as row
merge (country:Country { countryID: row.country_id})
on create set country.country = row.country;

load csv with headers from 'file:///customer.csv' as row
merge (customer:Customer { customerID: row.customer_id})
on create set customer.firstName = row.first_name
on create set customer.lastName = row.last_name
on create set customer.email = row.email;

```

```

load csv with headers from 'file:///film.csv' as row
merge (film:Film { filmID: row.film_id })
on create set film.title = row.title
on create set film.releaseYear = toInteger(row.release_year)
on create set film.rentalDuration = toInteger(row.rental_duration)
on create set film.rentalRate = toFloat(row.rental_rate)
on create set film.length = toInteger(row.length)
on create set film.rating = row.rating;

load csv with headers from 'file:///inventory.csv' as row
merge (inventory:Inventory { inventoryID: row.inventory_id });

load csv with headers from 'file:///language.csv' as row
merge (language:Language { languageID: row.language_id })
on create set language.name = row.name;

load csv with headers from 'file:///payment.csv' as row
merge (payment:Payment { paymentID: row.payment_id } )
on create set payment.amount = toFloat(row.amount)
on create set payment.paymentDate = row.payment_date;

load csv with headers from 'file:///rental.csv' as row
merge (rental:Rental { rentalID: row.rental_id } )
on create set rental.rentalDate = row.rental_date
on create set rental.returnDate = row.return_date;

load csv with headers from 'file:///staff.csv' as row
merge (staff:Staff { staffID: row.staff_id })
on create set staff.firstName = row.first_name
on create set staff.lastName = row.last_name
on create set staff.email = row.email;

load csv with headers from 'file:///store.csv' as row
merge (store:Store { storeID: row.store_id });
//
// Foreign keys
//
load csv with headers from 'file:///address.csv' as row
match (address:Address { addressID: row.address_id } )
match (city:City { cityID: row.city_id } )
merge (address)-[r:ADDRESS_CITY]->(city);

load csv with headers from 'file:///city.csv' as row
match (city:City { cityID: row.city_id } )
match (country:Country { countryID: row.country_id } )
merge (city)-[r:COUNTRY_OF_CITY]->(country);

load csv with headers from 'file:///customer.csv' as row
match (customer:Customer { customerID: row.customer_id } )
match (store:Store { storeID: row.store_id } )
match (address:Address { addressID: row.address_id } )
merge (customer)-[r1:CUSTOMER_STORE]->(store)
merge (customer)-[r2:CUSTOMER_ADDRESS]->(address);

load csv with headers from 'file:///film.csv' as row
match (language:Language { languageID: row.language_id } )
match (film:Film { filmID: row.film_id } )

```

```

merge (film)-[r:film_language]->(language);

load csv with headers from 'file:///inventory.csv' as row
match (inventory:Inventory { inventoryID: row.inventory_id} )
match (film:Film { filmID: row.film_id} )
match (store:Store { storeID: row.store_id} )
merge (store)-[r1:STORE_INVENTORY]->(inventory)
merge (film)-[r2:film_inventory]->(inventory);

load csv with headers from 'file:///payment.csv' as row
match (payment:Payment { paymentID: row.payment_id} )
match (customer:Customer { customerID: row.customer_id} )
match (staff:Staff { staffID: row.staff_id} )
match (rental:Rental { rentalID: row.rental_id} )
merge (payment)-[r1:PAYMENT_CUSTOMER]->(customer)
merge (payment)-[r2:PAYMENT_STAFF]->(staff)
merge (payment)-[r3:PAYMENT_RENTAL]->(rental);

load csv with headers from 'file:///rental.csv' as row
match (rental:Rental { rentalID: row.rental_id} )
match (inventory:Inventory { inventoryID: row.inventory_id} )
match (customer:Customer { customerID: row.customer_id} )
match (staff:Staff { staffID: row.staff_id} )
merge (rental)-[r1:RENTAL_INVENTORY]->(inventory)
merge (rental)-[r2:RENTAL_CUSTOMER]->(customer)
merge (rental)-[r3:RENTAL_STAFF]->(staff);

load csv with headers from 'file:///staff.csv' as row
match (staff:Staff { staffID: row.staff_id} )
match (address:Address { addressID: row.address_id} )
match (store:Store { storeID: row.store_id} )
merge (staff)-[r1:STAFF_ADDRESS]->(address)
merge (staff)-[r2:STAFF_STORE]->(store);

load csv with headers from 'file:///store.csv' as row
match (store:Store { storeID: row.store_id} )
match (staff:Staff { staffID: row.manager_staff_id} )
match (address:Address { addressID: row.address_id} )
merge (store)-[r1:STORE_MANAGER]->(staff)
merge (store)-[r2:STORE_ADDRESS]->(address);
// 
// Join tables for foreign keys
//
load csv with headers from 'file:///film_actor.csv' as row
match (actor:Actor { actorID: row.actor_id} )
match (film:Film { filmID: row.film_id} )
merge (actor)-[r:ACTS_IN]->(film);

load csv with headers from 'file:///film_category.csv' as row
match (film:Film { filmID: row.film_id} )
match (category:Category { categoryID: row.category_id} )
merge (film)-[r:film_category]->(category);

```

Importing the Pagila database takes about 10 minutes and will yield a graph that can be explored visually using Neo4J Browser using the following Cypher command that

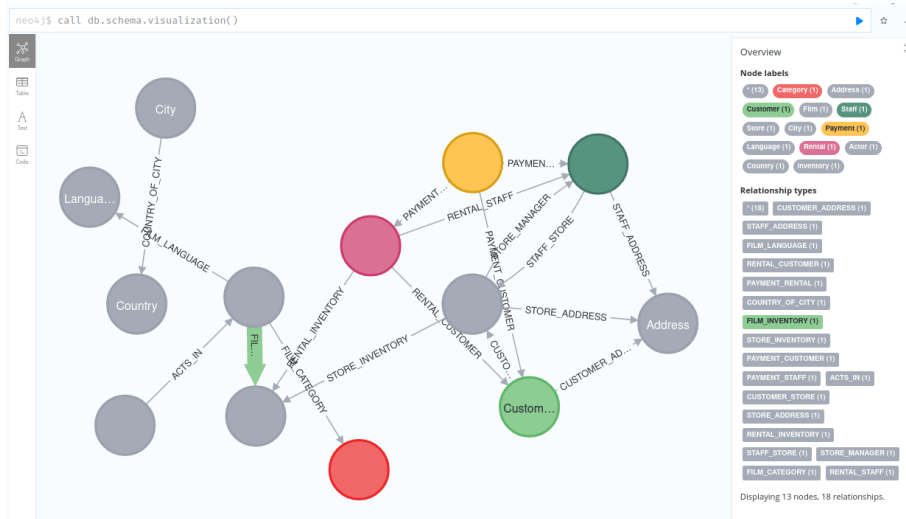


Figure 4.8: The Pagila database in Neo4J Browser

calls a built-in function for visualizing the database schema. A screen shot of the visual explorer is shown in Figure 4.8.

```
CALL db.schema.visualization()
```

When importing from files, or exporting to files, Neo4J Community Edition uses the the `/var/lib/neo4j/import/` directory on the server. Files to import must be placed in that directory, and exported files will be created there. Additionally, any scripts to be run by calling `CALL apoc.cypher.runFile()` must be located in that directory.

4.8 Graph Queries with Cypher

This section introduces the syntax of Cypher queries using example queries for the Pagila database as imported in the previous section.

Example: Find actors by last name, limit to 10.

```
MATCH (a:Actor)
RETURN a.firstName, a.lastName
ORDER BY a.lastName DESC
LIMIT 10;
```

The Cypher code above shows basic node label matching in the MATCH clause, returning a selection of node properties using the RETURN clause, ordering and limiting the result set using the ORDER BY and LIMIT clause, which are analogous to the SQL clauses with the same names.

Example: Find films whose title starts with a 'T' and that have a rental rate less than 3, sort by film title, limit to 10.

```
MATCH (f:Film {rating: "PG"})
WHERE (f.title STARTS WITH "T") AND (f.rentalRate < 3)
RETURN f.title, f.rating, f.rentalRate
ORDER BY f.title ASC LIMIT 10;
```

The Cypher code above introduces note matching on labels and properties and filtering using a WHERE clauses. Two conditions are combined using the AND word. Note that the matching on the rating property value of 'PG' could also have been incorporated into the WHERE clause, but not all WHERE clause conditions can always be moved to the node property specification in the MATCH clause and queries may be more readable when using a WHERE clause.

Example: Find rental datas and customer names of customers that live in India.

```
MATCH (r:Rental)
  - [:RENTAL_CUSTOMER] -> (c)
    - [:CUSTOMER_ADDRESS] -> ()
      - [:ADDRESS_CITY] -> ()
        - [:COUNTRY_OF_CITY] -> (ct {country: "India"})
RETURN c.firstName, c.lastName, r.rentalDate LIMIT 5
```

This example introduces matching of paths that contain multiple nodes and multiple relationships. In the above query, the types or labels of nodes and relationships are specified, but because no properties of the intermediate nodes or relationships are to be returned, they do not need to be bound to query variables.

Hands-On Exercise

Write a Cypher query to find all customers that have rented a film with rating "PG":

1. Explore the graph visually in Neo4J browser, note the relationship types (see Figure 4.9)
2. Consider the path from customer to film via rental and inventory
3. Design a pattern that starts with a customer node and ends with a film node
4. Define an appropriate WHERE clause of property restrictions in node patterns

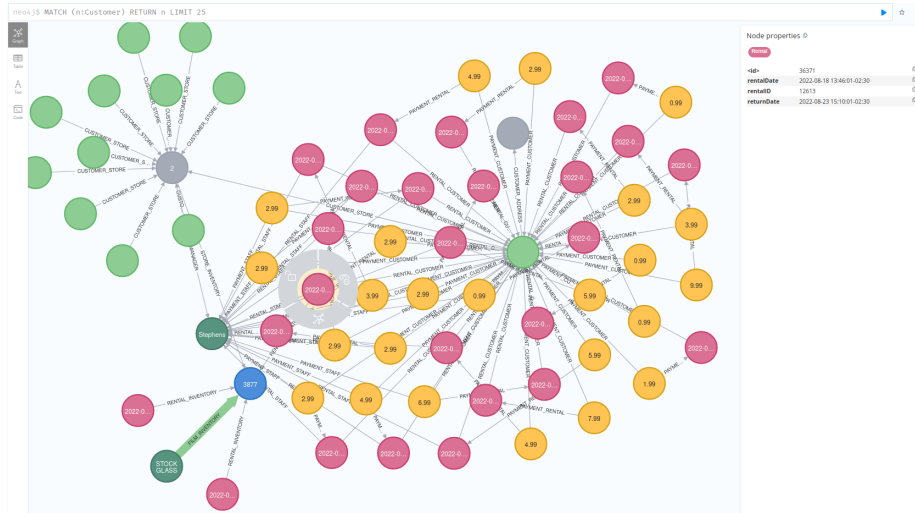


Figure 4.9: Exploring relationships among nodes in Neo4J Browser

Example: Find the mean and standard deviation of rental payments by country.

```

MATCH (p:Payment)
  - [:PAYMENT_RENTAL] -> (r:Rental)
  - [:RENTAL_CUSTOMER] -> (c)
  - [:CUSTOMER_ADDRESS] -> ()
  - [:ADDRESS_CITY] -> ()
  - [:COUNTRY_OF_CITY] -> (ct)
WITH ct,
     avg(p.amount) AS amountMean,
     stDev(p.amount) AS amountSD
RETURN ct.country, amountMean, amountSD
ORDER BY amountMean DESC LIMIT 5
  
```

This example introduces **aggregation**. In contrast to aggregation in SQL where grouping variables must be declared in the GROUP BY clause, grouping in Cypher is implicit and uses all non-aggregated variables. In the following example, the non-aggregated variable is `ct` (the country). The query also introduces the aggregation functions `avg()` and `stDev()` that compute the average and standard deviation, respectively. More information of aggregation functions can be found in the Neo4J documentation².

Example: Find the sets of last names of the movie cast, and the total number of actors.

²<https://neo4j.com/docs/cypher-manual/current/functions/aggregating/>

```
MATCH (a:Actor)-[:ACTS_IN]->(f:Film)
RETURN f.title,
       collect(a.lastName) AS cast,
       count(*) AS numActors;
```

This example introduces aggregation into collections (lists) using the `collect()` function. The query returns a list of actor last names as `cast`, together with the count of actors that act in each movie. Grouping happens implicitly for each variable not aggregated. In this example, that is the variable `f`, representing the film.

Example: Find the set of film titles by rental customer and the number of rentals.

```
MATCH (f:Film)-[:FILM_INVENTORY]-()
      -[:RENTAL_INVENTORY]->(r:Rental)
      -[:RENTAL_CUSTOMER]->(c:Customer)
RETURN c.lastName,
       collect(f.title) AS filmRentals,
       count(*) AS numRentals;
```

This example also uses aggregation with collection and a slightly more complex graph pattern in the `MATCH` clause³.

Example: Find the set of rental customers for each film and the rental count.

```
MATCH (f:Film)-[:FILM_INVENTORY]-()
      -[:RENTAL_INVENTORY]->(r:Rental)
      -[:RENTAL_CUSTOMER]->(c:Customer)
RETURN DISTINCT f.title,
       collect(c.lastName" "+left(c.firstName,1)+".") AS custNames,
       count(*) as rentalCount
```

This example introduces string functions and operators. Strings can be concatenated with the "+" operator. The function `left(., n)` returns the leftmost segment of `n` characters of the string. In contrast to the last query, here the collection creates a list of customers, grouped by films, rather than films, grouped by customer. The query also introduces the `DISTINCT` key word that limits the result set to unique values of a variable.

Example: Find the customers who rent films that are in inventory at multiple stores.

³From <https://neo4j.com/docs/getting-started/cypher-intro/results/>

```

MATCH (c:Customer)-[:RENTAL_CUSTOMER]-(r:Rental)
  -[:RENTAL_INVENTORY]-()
  -[:FILM_INVENTORY]-(f:Film)
WITH c, count{
  MATCH (f)-[:FILM_INVENTORY]-()
    -[:STORE_INVENTORY]-(s:Store)
  RETURN DISTINCT s.storeID } AS storeNum
WHERE storeNum > 1
RETURN DISTINCT
  c.lastName+" "+left(c.firstName,1)+"." AS custName,
  storeNum

```

This example introduces *sub-queries* and the `WITH` clause. The `WITH` clause introduces elements that will be passed to subsequent clauses. In this example, the result of the subquery within the `{ ... }` function is passed on in the variable `storeNum` in the "outer" query. This is then used in the `WHERE` clause of the outer query.

Example: Find Christian Akroyd's co-actors.

```

MATCH (a:Actor {firstName:"CHRISTIAN", lastName:"AKROYD"})
  -[:ACTS_IN]->(f:Film)-<[:ACTS_IN]-(coActors)
RETURN coActors.firstName+" "+coActors.lastName AS Name;

```

This query example emphasizes path matching from a given node. Note the second `ACTS_IN` relationship is traversed in reverse order, it's arrow points "left".

Example: Movies and actors up to 2 "hops" away from Christian Akroyd.

```

MATCH (a:Actor {firstName:"CHRISTIAN", lastName:"AKROYD"})
  -[:ACTS_IN*1..2]-(others:Actor)
RETURN distinct others;

```

This query introduces *quantified relationships*. In the example, the `ACTS_IN` relationship may be traversed between 1 and 2 two times on the way to other actor nodes. Note that no Film nodes or other relationships need to be specified here.

Example: The shortest path of an acts-in relationship between Christian Akroyd and Charlize Theron.

```

MATCH path=shortestPath(
  (a1:Actor {firstName:"CHRISTIAN", lastName:"AKROYD"})
  -[:ACTS_IN*]-(a2:Actor {firstName:"CHARLIZE", lastName:"THERON"}))
RETURN path;

```

This query introduces the use of *built-in functions*. In this case, the built-in function `shortestPath()` is a graph-theoretic function that computes the shortest path along `ACTS_IN` relationships between two specific nodes. Graph databases are particularly useful and efficient for queries on such graph-theoretic functions, which are very difficult to express in SQL.

Example: Find actors that Christian Akroyd hasn't yet worked with, but his co-actors have. Extend Christian Akroyd's co-actors, to find co-co-actors who haven't worked with him.

```

MATCH (a1:Actor {firstName:"CHRISTIAN", lastName:"AKROYD"})
      -[:ACTS_IN]->(m)<-[ :ACTS_IN]-(coActors),
      (coActors)-[:ACTS_IN]->(m2)<-[ :ACTS_IN]-(cocoActors)
WHERE NOT (a1)-[:ACTS_IN]->()<-[ :ACTS_IN]-(cocoActors)
      AND a1 <> cocoActors
RETURN cocoActors.firstName+" "+
       cocoActors.lastName AS Recommended,
       count(*) AS Strength
ORDER BY Strength DESC
    
```

This query example introduces the use of multiple patterns in the `MATCH` clause that are separated by commas and are related in the sense that variables in one can be used in the other and refer to the same node or relationship. The two patterns in the `MATCH` clause are connected through the shared variable `coActors`. Note also that traversal direction of the various relationships. Finally, this example also introduces the use of patterns in the `WHERE` clause, allowing more complex filters on the results. The patterns in the `WHERE` clause are also logically related to the patterns in the `MATCH` clause, in this example they share the variable `cocoActors`.

Example: Find someone who can introduce Christian Akroyd to Susan Davis.

```

MATCH (a1:Actor {firstName:"CHRISTIAN", lastName:"AKROYD"})
      -[:ACTS_IN]->(m)<-[ :ACTS_IN]-(coActors),
      (coActors)-[:ACTS_IN]->(m2)
      <-[ :ACTS_IN]-(a2:Actor {firstName:"SUSAN", lastName:"DAVIS"})
RETURN a1, m, coActors, m2, a2
    
```

The example is similar to the one above with its use of multiple path patterns in the `MATCH` clause. The query finds common co-actors of two named actors. Note the traversal directions of the relationships.

Hands-On Exercises

The following hands-on exercises are designed to familiarize you with the Cypher language and use the Pagila database.

1. Are there two customers that have the same address?
2. Which customers have rented the same set of films?
3. Find all films with a single actor
4. Calculate the rental revenue per customer. Who are the top 5? Bottom 5?
5. Calculate the rental counts for each country of customer. Are there countries with no rentals?
6. Create a graph that represents a product hierarchy.

4.9 Review Questions

1. What is a graph database, and how does it differ from traditional relational databases?
2. Describe the different data models used in NoSQL databases. How does the graph model specifically cater to certain types of data and applications?
3. Explain how data is represented in a graph database. What are nodes and relationships?
4. List and explain the key benefits of using graph databases over traditional relational databases.
5. How do graph databases handle relationships differently, and why is this advantageous for certain applications?
6. Give examples of specific industries or applications where graph databases are particularly useful. Explain why a graph database is chosen over other types of databases in these scenarios.
7. What are some of the prominent query languages used with graph databases? Briefly describe their unique features.
8. How does Cypher, the query language for Neo4j, compare to SQL in terms of syntax and capabilities?
9. Discuss the characteristics of Cypher as a query language. How does it enable efficient querying and manipulation of graph data?
10. Reflect on a scenario or a problem where you think a graph database would be more effective than a traditional relational database. Explain your reasoning.
11. Describe what a node represents in Neo4j and how it is represented in Cypher syntax.
12. Explain how properties are associated with nodes in Neo4j. Give an example using Cypher syntax.
13. Discuss the significance of relationship directionality in Neo4j. What is the difference between directed and undirected relationships in querying?
14. Define what a 'pattern' is in Cypher and its role in querying the graph database.
15. Provide an example of a simple Cypher pattern and explain what it matches in

the graph.

16. Differentiate between the ‘CREATE’ and ‘MERGE’ statements in Cypher. Under what circumstances would you use each?
17. Give an example of how to create a node with multiple labels and properties using Cypher.
18. How would you create a relationship between two nodes, including setting properties on the relationship?
19. Explain the difference between modeling data as a property of a node versus as a separate node connected by a relationship. Give an example to illustrate your point.
20. In the context of Neo4j, why might it be more efficient to model certain data as relationships between nodes rather than as properties of a single node? Provide an example where this is the case.
21. Given a graph model where movie genres are modeled as properties of a movie node, what are the limitations of this approach when trying to find movies with shared genres?
22. Describe the process of translating a relational data model into a graph model in Neo4j.

Chapter 5

Introduction to Data Management with R

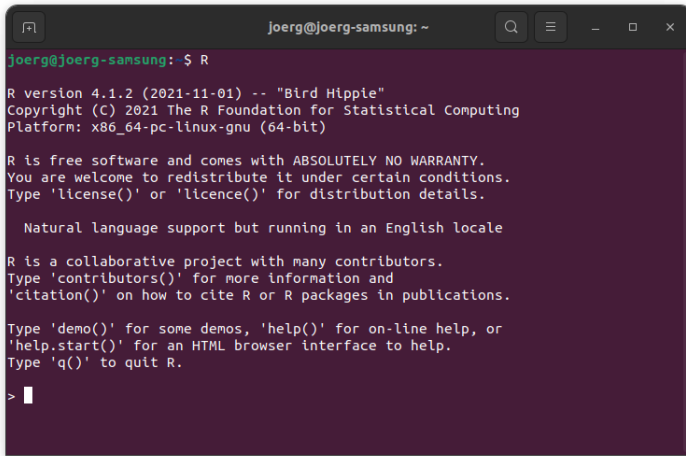
5.1 Introduction

R is a highly acclaimed statistical software and programming language known for its robust capabilities in data analysis, visualization, and statistical computing. It was conceived in the early 1990s by Ross Ihaka and Robert Gentleman at the University of Auckland, New Zealand. Drawing inspiration from the S language developed at Bell Laboratories, R was designed to be a powerful and flexible tool for data analysis and statistical modeling.

One of the key advantages of R is its open-source nature, making it freely available to users worldwide. This accessibility has fostered a vibrant community of users and developers, continuously enhancing its functionality through comprehensive packages and extensions. The Comprehensive R Archive Network (CRAN), a repository of these packages, is a testament to R's extensible architecture, offering tools for a myriad of data analysis tasks.

R's popularity stems not only from its wide range of statistical techniques, including linear and nonlinear modeling, time-series analysis, classification, clustering, and others, but also from its exceptional capabilities in data visualization. The software provides an integrated suite of tools for data manipulation, calculation, and graphical display, making it an invaluable asset for statisticians, researchers, and data scientists.

Moreover, R's programming language aspect allows for automation and customization in data analysis, which is highly beneficial for complex and repetitive tasks. Its compatibility with various data formats and integration with other programming languages and tools further enhances its versatility.



```
joerg@joerg-samsung: ~
R version 4.1.2 (2021-11-01) -- "Bird Hippie"
Copyright (C) 2021 The R Foundation for Statistical Computing
Platform: x86_64-pc-linux-gnu (64-bit)

R is free software and comes with ABSOLUTELY NO WARRANTY.
You are welcome to redistribute it under certain conditions.
Type 'license()' or 'licence()' for distribution details.

Natural language support but running in an English locale

R is a collaborative project with many contributors.
Type 'contributors()' for more information and
'citation()' on how to cite R or R packages in publications.

Type 'demo()' for some demos, 'help()' for on-line help, or
'help.start()' for an HTML browser interface to help.
Type 'q()' to quit R.

> ■
```

Figure 5.1: The R command line interface

5.2 Using R

R is a command-line oriented software, that is, users type commands to perform calculations or call functions of R packages. A sequence of R commands can be assembled in a *script file*, so that they may be re-run when necessary. The advantage of this type of software over one with a graphical user interface is in the repeatability and replicability of the work. Ideally, data analysts will assemble an R script file for their entire data analysis, from raw data sets to finished statistical analyses and visualizations, so that all details of the analysis are available for replication and evaluation.

The R system can be launched simply by invoking the `R` command from the terminal window, as shown in Figure 5.1. R will display its version information and prompt for command entry with a `>` prompt.

To install R on Microsoft Windows or on MacOS, download the installation files from CRAN (Comprehensive R Archive Network) at <https://cran.r-project.org> and follow the instructions. R on Microsoft Windows and R on MacOS will show their command prompts inside a window but otherwise function similarly to R on Ubuntu that is installed in the course virtual machine.

Tip: A good, easy, and comprehensive introduction to R can be found here:
<https://cran.r-project.org/doc/manuals/r-release/R-intro.pdf>

Tips for working efficiently with R: To make using R more efficient, consider doing the following:

- Use the up-arrow key to retrieve earlier commands.
- The `history()` function shows your command history.
- Use a notepad app to assemble your commands, then copy/paste to R.
- Use a notepad app for your results, copy/paste from R.
- The Ubuntu terminal window uses SHIFT-CTRL-X, SHIFT-CTRL-C, SHIFT-CTRL-V for cut/copy/paste.
- Use multiple terminal and R windows (e.g. one for executing commands, one for reading help documentation or for listing files).
- Don't update packages in the middle of a project.
- Ensure you have a *repeatable, automatable script* for your entire data analysis at the end of a project.

5.3 R Basics

The most basic way to use R is to simply use it as a calculator, as shown in the following R code example. Type "1+1" at the ">" prompt, then press the RETURN key to execute the statement. R will respond on the following line with the result:

```
> 1+1
[1] 2
```

A *variable* in R is a named storage space for numbers, characters, strings, and other data elements. Traditionally, values are assigned to variables using the `<-` operator, but one may also use the more "normal" assignment operator `=`. Using the `<-` assignment operator helps to clearly distinguish assignment from equality testing, which uses `==`.

The following R code example introduces the R function called `print()` that does as its name suggests. Most data types and data structures that can be assigned to variables have a useful print function associated to them, so that on the interactive R command line you can simply type their name to get their value. In interactive mode, R calls the `print()` function automatically, in an R script that you execute from file, you will have to explicitly use the `print` function.

```
> a <- 3
> b <- 2
> print(a * b)
[1] 6
> a
[1] 3
```

A common structured data type in R is a *vector*. A vector in R contains elements of

the same data type and is ordered. When assigning elements of different datatypes to a vector, R will coerce the types of all elements to a common datatype.

```
> v <- c(1, 'a', TRUE)
> v
[1] "1"     "a"     "TRUE"
> v <- c(1, 2, 3, 4)
> v*3
[1] 3 6 9 12
```

Note that R automatically determined that multiplication with a scalar is an element-wise operation and applies it to each element of the vector.

Useful functions to create vectors are the sequence function `seq()`, which accepts the lower and upper limit and a step size as parameter, and the repetition function `rep()` which repeats its first argument the number of times specified by its second argument.

```
> s <- seq(0, 6, by=.5)
> print(s)
> r <- rep(3.5, 5)
> print(r)
```

R provides useful functions for numerical vectors, to find their length, their maximum and minimum value, the square root of their values, as well as the variance and standard deviation of the elements. Note that R automatically determines whether functions are applied to the whole vector, like `var()` or `sd()`, or whether functions are applied element-wise to each element, like `sqrt()`. Vector concatenation, using the `c()` function, automatically "flattens" the vectors.

```
> length(v)
> max(v)
> min(v)
> sqrt(vv)
> var(v)
> sd(v)
> vv <- c(v, c(7, 8, 9), v)
> print(vv)
```

The most common way to select elements from vectors is by *indexing* with a boolean vector. In the following example, the expression `vv < 5` yields a vector of boolean values. Indexing the variable `vv` with that vector determines which elements of `vv` to select.

```
> vv < 5
> vv[vv < 5]
> vv[vv < 5] <- vv[vv < 5] + 5
```

Vectors can also be indexed numerically, selecting elements by their position. R allows you to specify a sequence using the `:` operator and exclusion of elements using `-`. The first line in the following example selects elements at positions 3 through 7, the second line selects elements *except* those at positions 3 through 7.

```
> vv[3:7]
> vv[-(3:7)]
```

Important:

- R begins indexing positions with 1, while other programming languages begin at 0.

Tip:

- The boolean constants `TRUE` and `FALSE` can be abbreviated by `T` and `F`

R also has special symbols to denote infinity (`inf`) and results that are not a number (`NaN`):

```
> 2 / 0
[1] Inf
> 0 / 0
[1] NaN
```

Importantly, `NaN` is *not* the same as a missing value, which is denoted by `NA`, as in the following R code example. The `is.na()` function can be used to identify and index `NA` and then filter them. Any `NA` typically yields an `NA` when an aggregate function is applied. Many functions offer an option to remove `NA` values prior to applying them, as shown for the `sum()` function in the following R code block.

```
> v[3] <- NA
> v*3
[1] 3 6 NA 12
> is.na(v)
[1] FALSE FALSE TRUE FALSE
> sum(v)
[1] NA
> sum(v, na.rm=TRUE)
[1] 7
```

The boolean logical *and* and *or* are represented by the operators `&` and `|` shown in the R code block below.

```
> TRUE & FALSE
FALSE
> TRUE | FALSE
TRUE
```

Character strings in R are enclosed in single or double quotes (but not mixed quotes!). Two useful functions are `paste()` which pastes its arguments together with an optional separator between them and returns a characters string, and the `strsplit()` function which accepts a string (or vector of strings) to split, and a separator character that identifies where to split the string. It returns a list of vectors of strings.

```
> label1 = 'I Love R'
> label2 = 'and BUSI 4760'
> paste(label1, label2, sep=' ')
> strsplit('Hello World! My first string', ' ')
```

Because you can assign arbitrary values to variables in R, R provides functions to test the value type and to change or coerce the value type. A *factor* data type in R represents categorical data, encoded as different character strings or different numbers. Categorical data is treated different from numerical or character string data in many statistical analyses.

```
> is.numeric(vv)
> is.integer(vv)
> mode(vv)
> as.character(vv)
> is.character(as.character(vv))
> as.factor(as.character(vv))
> levels(as.factor(as.character(vv)))
```

Important string functions are `grep()`, which checks whether strings contain a substring that matches a regular expression, and `agrep()`, which calculates the Levenshtein distance between a regular expression and a set of strings. The Levenshtein distance is defined as the sum of insertions, deletions, and substitutions of characters to transform one string into another. The first use of `grep()` in the following R code block matches a phone number, the second use of `grep()` matches a Canadian postal code, while the last two examples of `grep()` and `agrep()` exemplify the difference between exact matching with `grep()` and approximate matching with `agrep()`.

```
> grep('^( [0-9]{3})[ -]?[0-9]{3}[ -]?[0-9]{4}$',
      c('709 864 5000', 'abc def 9999', '709-865-5000'))
[1] 1 3
> grep(' [A-V] [0-9] [A-V] [0-9] [A-V] [0-9] ',
      c('A0P 1L0', '0AB L2K', 'A0X 1Z0'))
[1] 1
> grep('apple', c('apricot', 'banana', 'grape', 'pineapple'))
[1] 4
> agrep('apple',
         c('apricot', 'banana', 'grape', 'pineapple'),
         max.distance=3)
[1] 1 3 4
```

5.4 The R Environment

The collection of variables, functions and libraries that exists in R at any one time is called the R *workspace*. R provides many functions to manipulate objects in its workspace, among them `ls()` and `rm()`, named after their Unix bash shell equivalents. The following R code illustrates the use of these functions. Results may vary depending on what variables have been created prior to these commands.

```
> ls()
[1] "a"          "b"          "v"
> rm(v)
> ls()
[1] "a"          "b"
```

R comes with a built-in user manual that one can access with the `help()` function or simply the `? operator`. Help is available on any function in R, as shown in the following example. For added convenience, R provides a web browser interface to its help pages that is started by `help.start()`.

```
> help()
> help(lm)
> ?lm
> ??lm
> help.start()
```

R has a working directory where it reads and writes files from and to. On Ubuntu Linux, this is the directory from which the `r` command was issued. R provides functions to get the working directory, to set (change) it, and to list the files in the working directory:

```
> getwd()
[1] "/home/busi4720"
> setwd('DataSets')
> getwd()
[1] "/home/busi4720/DataSets"
> list.files()
```

Tip: It is often more convenient to change the working directory in the terminal, prior to invoking `r`.

A collection of related functions is called a *library* in R. While some libraries come with the base R system, other packages will need to be downloaded and installed. The CRAN (comprehensive R archive network) provides libraries in convenient form. To install packages from CRAN, use the `install.packages()` which accepts the name (or a vector of names) of packages to install from CRAN. On some systems, R may prompt the user from which CRAN location to install packages. Normally, there is little difference other than download speed.

Installed libraries can be attached to the R workspace with the `library()` function. The `library()` function with an argument attaches the specified package and makes its functions and data sets available for use. The `library()` function without any arguments shows which libraries are installed. The `search()` function shows which packages are currently attached to the workspace. Finally, `installed.packages()` provides details of all installed packages.

```
> search()
> library(matrixcalc)
> search()
> library()
> install.packages('lavaan')
> library()
> installed.packages()
```

It is sometimes useful to assemble a set of related R commands in a script file. As noted earlier, script files are useful to improve the replicability of the data analysis. The `source()` function will read and execute a file containing R commands. As noted earlier, in a script file, you will need to use the `print()` function to print the values of variables.

```
> source('MyFirstScript.R')
```

Finally, the `quit()` function ends an R session. When using `quit()` without arguments, R will ask whether to save the workspace image. R stores its *workspace* in each

directory in a file called ".RData" and will read it when restarted from that directory. R also stores its *command history* in each directory in a file called ".Rhistory" and will read it when restarted from that directory.

```
> quit()
```

5.5 Arrays, Matrices, Lists, and DataFrames

R *arrays* are multi-dimensional objects that can hold any primitive data type, usually numerical. A *matrix* is simply a two-dimensional array. The following example shows how indexing generalizes from vectors to matrices and arrays simply by indexing each dimension with the same syntax as used for vectors. The `array()` creates multi-dimensional arrays from existing data, the `dim()` function returns the number of dimensions of an array.

A few important things to note in the following R code block example:

- Initially, the array is created from a range of numbers between 1 and 20, and the `dim` argument specifies the dimensionality.
- A dimension need not be subsetted or indexed, as in `a[, 2]` or `a[, 2:4]` which do not subset the first dimension
- Reversing the index reverses the result that is returned, as in `a[3:1, 2:4]` which reverses the indexing of the first dimension.
- An array with two columns is interpreted as a set of indexes, as in `a[i] <- 0`

```
> a <- array(1:20, dim=c(4,5))
> a
     [,1] [,2] [,3] [,4] [,5]
[1,]    1    5    9   13   17
[2,]    2    6   10   14   18
[3,]    3    7   11   15   19
[4,]    4    8   12   16   20
> dim(a)
[1] 4 5
> a[,2]
> a[,2:4]
> a[3,2:4]
> a[3:1,2:4]
> i <- array(c(1:3,3:1), dim=c(3,2))
> a[i] <- 0
> a
```

Constructing a *matrix* with the `matrix()` function is similar to constructing an array, but instead of providing the dimensionality with `dim`, one must provide the number of rows or columns (`nrow` or `ncol`) and how to fill the matrix from the elements provided

using the `byrow` argument. The `t` function returns the transpose of a matrix, that is, it reverses rows and columns. Binding two matrices together by columns with `cbind()` or by rows with `rbind()` requires compatible dimensions.

```
> b <- matrix(20:1, nrow=5, byrow=T)
> b
 [,1] [,2] [,3] [,4]
[1,] 20   19   18   17
[2,] 16   15   14   13
[3,] 12   11   10   9
[4,] 8    7    6    5
[5,] 4    3    2    1
> is.matrix(b)
> is.matrix(a)
> t(b)
> cbind(a, t(b))
> rbind(t(a), b)
```

A *list* in R is an ordered collection of elements that, in contrast to vectors, *may be of different types*. Lists are created using the `list()` function. Note the difference in selecting elements: The `[[]]` operator returns the element at that position in the list, whereas the `[]` operator contains a list that contains the element at that position in the list.

```
> l <- list('a', 3, 'b', 2, TRUE)
> l[[2]]
> l[2]
> is.list(l)
> is.list(l[[2]])
> is.list(l[2])
> as.list(vv)
```

A *data frames* are the most widely used data structure for data analytics and statistics in basic R. It is essentially a table with a set of columns whose elements are of the same type. Columns are named and columns can be selected using the `$` symbol. Useful functions on data frames are `summary()`, `head()` and `tail()`. The following R code block creates a variable `x` as a vector of 50 normally distributed random values using the `rnorm()` function. The variable `y` is created from vector `x` and additional normally distributed random variables. The two are then combined into a data frame. The `colnames()` function can retrieve the column names, but can also be used to change/update the column names. The `nrow()` and `ncol()` functions return the number of rows and columns, `head()` and `tail()` return the first few or last few rows, and `cov()` is an example of a statistical function that returns the covariance matrix of all columns in the data frame.

```
> x <- rnorm(50)
> y <- 2*x + rnorm(50)
> data <- data.frame(x, y)
> colnames(data)
> colnames(data) <- c('Pred', 'Crit')
> nrow(data)
> ncol(data)
> data$Pred
> summary(data)
> head(data)
> tail(data)
> cov(data)
```

Data frames may be written to CSV files and read from CSV files, as shown in the following R code block. Both functions, `write.csv()` and `read.csv()` have a range of options for reading/writing files with or without header lines, different separators, for skipping rows, different decimal points, whitespace stripping, etc. Consult the R built-in help system for details.

```
> write.csv(data, 'data.csv', row.names=FALSE)
> new.data <- read.csv('data.csv')
> colnames(new.data)
```

5.6 Tidyverse

The Tidyverse is a collection of R libraries designed for data science that share an underlying design philosophy, grammar, and data structures. Developed by Hadley Wickham and others, the Tidyverse libraries are built to work together seamlessly, making data science tasks more straightforward and intuitive. At the core of Tidyverse's philosophy is the concept of "tidy data," which arranges data in a structured way that simplifies analysis. This structure involves organizing data into rows and columns where each variable is a column, each observation is a row, and each type of observational unit forms a table.

Key libraries in the Tidyverse include `ggplot2` for data visualization, `dplyr` for data manipulation, `tidyr` for data tidying, `readr` for reading data, `purrr` for functional programming, and `tibble` for providing a better version of a table data structure. In particular, `ggplot2` allows for complex and aesthetically pleasing visualizations using a layered grammar of graphics (hence the name), while `dplyr` provides a set of tools for efficiently manipulating datasets, such as filtering rows, selecting columns, and aggregating data. `tidyverse` helps in transforming messy data into a tidy format, making it easier to analyze and visualize. Table 5.1 contains a summary of the libraries.

The Tidyverse also emphasizes readability and expressiveness in code, which not only makes data analysis easier to write but also easier to read and understand. It has become

dplyr	Manipulate data
forcats	Work with categorical variables (factors)
ggplot2	Grammar of Graphics
lubridate	Date and time parsing and arithmetic
purrr	Functional programming
readr	Read files in various formats
stringr	Work with character strings
tibble	A tibble is better than a table
tidyverse	Make data tidy

Table 5.1: Tidyverse packages for R

```
> library(tidyverse)
— Attaching core tidyverse packages ————— tidyverse 2.0.0 —
✓ dplyr    1.1.3   ✓ readr    2.1.4
✓ forcats  1.0.0   ✓ stringr  1.5.0
✓ ggplot2  3.4.4   ✓ tibble   3.2.1
✓ lubridate 1.9.3  ✓ tidyverse 1.3.0
✓ purrr   1.0.2
— Conflicts ————— tidyverse_conflicts() —
✖ dplyr::filter() masks stats::filter()
✖ dplyr::lag()    masks stats::lag()
i Use the conflicted package to force all conflicts to become errors
> █
```

Figure 5.2: Attaching the tidyverse packages in R

a popular choice among data scientists and statisticians for its ease of use, efficiency, and the cohesive way it handles data analysis tasks. The integration of these packages under the Tidyverse umbrella simplifies the process of data manipulation, exploration, and visualization, greatly enhancing the productivity and effectiveness of data analysis in R.

Tip: An introduction to data science with the Tidyverse packages, directly from their authors, can be found here: <https://r4ds.hadley.nz/>

Loading and attaching the `tidyverse` library in R, using the `library(tidyverse)` function, loads all the associated core packages, as shown in Figure 5.2.

This section can give only a very brief outline of the capabilities of the tidyverse packages. The extensive documentation and various "cheat sheets"¹ provide additional details. This section focuses on the use of `dplyr` to analyze data from a set of CSV files representing the data of the Pagila database. The Pagila database² is a demonstration

¹<https://posit.co/resources/cheatsheets/>

²<https://github.com/devrimgunduz/pagila>,

<https://github.com/devrimgunduz/pagila/blob/master/LICENSE.txt>

database originally developed for teaching and development of the MySQL RDBMS under the name Sakila³. Pagila is designed as a sample database to illustrate database concepts and is based on a fictional DVD rental store. It originally consists of several tables organized into categories like film and actor information, customer data, store inventory, and rental transactions. For this section, the Pagila data was summarized in a few related CSV files.

When reading CSV files with `readr`, the data is stored in a *tibble*, not a data frame. A tibble provides a number of extensions and convenience operations that make it significantly more capable than a data frame. When using data frames with `dplyr`, they are automatically converted to tibbles.

The following R code reads a CSV file using the `read_csv()` function and prints the first few lines and a summary. The output looks slightly different than that for data frames, but accomplishes essentially the same things.

```
rentals <- read_csv('rentals.csv')
head(rentals)
summary(rentals)
```

Attaching a tibble or data frame with `attach()` means that its columns become variables in the R workspace and need not be selected from the tibble (or data frame) using the `$` operator. The following R code block transforms the data read from the CSV file into the `rentals` tibble into appropriate data types, then detaches the tibble and prints a summary.

```
# Fix the column datatypes:
attach(rentals)
rating <- as.factor(rating)
language <- as.factor(language)
customer_address <- as.integer(customer_address)
customer_store <- as.integer(customer_store)
rental_staff <- as.integer(rental_staff)
payment_staff <- as.integer(payment_staff)
rental_duration <- as.integer(rental_duration)
detach(rentals)
summary(rentals)
```

The tidyverse libraries make extensive use of the *pipe* operator in R. The pipe operator allows chaining of function calls and plugs the result of one function as the first argument into the next function. Originally, tidyverse used the `%>%` pipe operator from the `magrittr` library, but can now also be used with the new (since R version 4.1), native R pipe operator `|>`. For simple usage, the two behave identically and can be intermixed.

³<https://dev.mysql.com/doc/sakila/en/>,
<https://dev.mysql.com/doc/sakila/en/sakila-license.html>

The following R code block demonstrates a simple sequence of data manipulation operations using functions from the dplyr library. It begins with the data tibble which is piped into the first function. The outputs of each function are piped into the following function, ending with `print()`. Note that print output can also be piped into other functions, allowing printing of intermediate results.

```
rentals |>
  filter(if_any(everything(), is.na)) |>
  select(last_name, rental_date, return_date, title, amount) |>
  print(n=Inf, width=Inf)
```

- The `filter(if_any(everything(), is.na))` function is the first in the pipeline. It filters rows in the rentals data frame based on the presence of NA (missing) values. The `if_any()` function checks each column (indicated by `everything()`) for NA values. The filter function then retains only those rows that have at least one NA value in any column.
- Next, the `select()` function specifies the columns to retain in the resulting data frame. It narrows down the data frame to include only the `last_name`, `rental_date`, `return_date`, `title`, and `amount` columns. This step reduces the dataset to focus on these key variables, for easier analysis and reporting.
- Finally, the `print(n=Inf, width=Inf)` function displays the output. The `n=Inf` argument tells R to print all rows of the data frame, instead of just the first few rows as is the default behavior. Similarly, `width=Inf` ensures that all columns are printed without any being truncated, which is useful for wide tibbles or data frames.

In summary, this R code example is used to filter a rentals table for rows containing missing values in any column, and then to select and print specific columns of interest. This kind of operation is typical in data cleaning and exploratory data analysis processes. The result shows that some films have not been rented (i.e. there is no rental date for them), and some rentals have not been returned (i.e. there is no return date for them).

The following paragraphs show examples of further data analysis with Tidyverse, introducing additional dplyr functions and their use. Dplyr functions are intended to mirror the SQL queries from the earlier chapter on relational databases. The main dplyr "verbs" used in the examples are summarized in Table 5.2.

Example: Find all films and the actors that appeared in them, ordered by film category and year, for those films that are rated PG.

full_join	Joins tibbles (also outer_join, left_join, inner_join, right_join)
filter	filters by row
select	selects columns to retain
mutate	creates new columns
rename	renames columns
distinct	finds unique values
group_by	groups data
nest	nests data, tibbles in tibbles
arrange	sorts data rows
relocate	moves data columns
print	prints a tibble

Table 5.2: Important dplyr functions

```

actors <- read_csv('actors.categories.csv')

rentals |>
  full_join(actors, by='title',
             suffix=c('_customer', '_actor'),
             relationship='many-to-many') |>
  filter(rating == 'PG') |>
  mutate(actor =
    paste(last_name_actor, ' ', first_name_actor, sep='')) |>
  rename(year=release_year) |>
  select(actor, title, category, year) |>
  distinct(actor, title, category, year) |>
  group_by(category, year, title) |>
  nest() |>
  arrange(category, year, title) |>
  relocate(category, year, title) |>
  print(n=Inf, width=Inf)

```

This R code processes the rentals tibble and the actors tibbles through a sequence of functions in a pipeline.

- The `read_csv()` function is used to read data from a CSV file named "actors.categories.csv" into an R data frame called "actors".
- The "rentals()" data frame is combined with the "actors" data frame using a full join. The join is performed on the "title" column common to both data frames. The `suffix` argument adds distinct suffixes to column names from each data frame to avoid name clashes. The `relationship='many-to-many'` indicates the nature of the join.

- The `filter()` on the combined data retains only rows where the "rating" column has the value "PG".
- The `mutate()` function is used to create a new column named "actor", which concatenates the actor's last name and first name, separated by a comma and a space.
- The "release_year" column is renamed to 'year' using the `rename` function.
- The `select()` function is used to narrow down the data frame to only the columns "actor", "title", "category", and "year".
- Following this, the `distinct()` function ensures that only unique rows are retained, removing any duplicates.
- The data is grouped by "category", "year", and "title", and then `nest` is used to create a nested data frame, i.e. a dataframe where the actors for each group are in a list-valued columns.
- the `arrange()` sorts the data frame by "category", "year", and "title", while `relocate` moves these columns to the front of the data frame for easier viewing.
- Finally, the entire data frame is printed with all rows (`n=Inf`) and without truncating any columns (`width=Inf`).

Example: Find the most popular actors in the rentals in each city.

This R code block below involves combining multiple data frames and then manipulating and summarizing the data. It builds on the rental and actor tibbles from the previous example and includes address information.

```
addresses <- read_csv('addresses.csv')
addresses$phone <- as.character(addresses$phone)

full_data <-
rentals |>
  inner_join(addresses, by=c('customer_address'='address_id')) |>
  inner_join(actors, by='title',
  suffix=c('_customer', '_actor'),
  relationship='many-to-many')

full_data |>
  mutate(actor =
  paste(last_name_actor, ', ', first_name_actor, sep='')) |>
  group_by(city, actor) |>
  summarize(count=n()) |>
  mutate(ranking = min_rank(desc(count))) |>
  filter(ranking < 4) |>
  arrange(city, ranking, actor) |>
  print(n=25)
```

- The analysis starts by reading a CSV file containing addresses into a data frame.
- An inner join is first performed between these two data frames, matching them on a specified key.
- This is followed by another inner join with an ‘actors’ data frame. This second join involves a many-to-many relationship and adds suffixes to overlapping column names to distinguish them.
- With the full data set, a new column is created by concatenating the first and last names of actors, forming complete names.
- The data is then grouped by city and actor.
- A new summary column is created that counts the number of occurrences (records) for each group.
- To create rankings, a new column is added that ranks the groups based on the count in descending order. The `min_rank()` function allows ties in the ranking, use `rank()` to break ties with gaps in ranking or `dense_rank()` to break ties with no gaps in ranking.
- The data is then filtered to include only those records with a ranking less than 4, focusing on the top three ranks for each group.
- Finally, the data is sorted by city, ranking, and actor and then printed.

Example: Find the customers who spend the most on rentals, with their phone numbers and cities, and the number of rentals with the highest total rental payments for each category grouped by rental duration.

```
full_data |>
  mutate(customer= paste(first_name_customer, last_name_customer)) |>
  select(customer, amount, rental_duration, category, phone, city) |>
  group_by(category, rental_duration, customer) |>
  mutate(payments=sum(amount), num_rentals=n()) |>
  select(-amount) |>
  group_by(category, rental_duration) |>
  mutate(ranking = min_rank(desc(payments))) |>
  slice(which.min(ranking)) |>
  print(n=Inf, width=Inf)
```

By now, it should be clear what the functions in the analysis pipelines accomplish. However, a few interesting things to note. First, there is no `summarize()` function because `summarize()` omits all non-grouped columns, but the example requires phone numbers and cities of customers. Either these would need to be included somehow in the `summarize()` function, or as is done in this R code, summary columns are created with `mutate()`. Second, note the “negative” argument to the `select()` function, which is used to remove the “amount” column. Third, the pipeline uses multiple `group_by()` statements with different aggregate functions (`sum()`, `n()`,

`min_rank()`) for the different groups. Finally, the R code uses `slice()` to select the rows with the smallest ranks.

Example: Get the total rental revenue, number of rentals, and the mean and standard deviation of the rental amounts for each country.

```
full_data |>
  group_by(country) |>
  summarize(revenue=sum(amount),
            numrentals=n(),
            mean_amount=mean(amount),
            sd_amount=sd(amount)) |>
  arrange(desc(mean_amount),
          desc(revenue)) |>
  print(n=Inf, width=Inf)
```

The R code for this query demonstrates a number different aggregate summary functions, `sum()`, `n()`, `mean()` and `sd()` (standard deviation). It also shows how to use the `desc()` function to arrange or sort data in decreasing order.

Example: Get the top 5 and the bottom 5 grossing customers for each quarter.

```
full_data |>
  mutate(customer=paste(first_name_customer, last_name_customer)) |>
  mutate(q=as.character(quarter(rental_date, with_year=T))) |>
  select(customer, q, amount, rental_date) |>
  group_by(q, customer) |>
  mutate(payments=sum(amount)) |>
  select(-amount) |>
  distinct(customer, q, payments) |>
  group_by(q) |>
  mutate(rank_top = min_rank(desc(payments))) |>
  mutate(rank_bot = min_rank(payments)) |>
  filter(rank_top < 6 | rank_bot < 6) |>
  arrange(q, desc(payments)) |>
  relocate(q, customer, payments, rank_top, rank_bot) |>
  print(n=Inf, width=Inf)
```

The code for this query again does not use a `summarize()` function. It also shows the use of the `quarter()` function from the "lubridate" library. The lubridate library contains a large range of date and time related functions. Two ranking columns are created using the `mutate()` and `min_rank()` functions, once in descending order to get the top ranks, and again in ascending order to get the bottom ranks. The code uses `filter()` instead of `slice()` to select the top and bottom 5 ranks, uses `arrange()` to sort the data, and then uses `relocate()` to re-arrange the order of columns prior to printing.

Example: Find the set of film titles by rental customer and the total number rentals for each customer.

```
full_data |>
  mutate(customer=paste(first_name_customer, last_name_customer)) |>
  select(customer, title) |>
  nest(titles=title) |>
  rowwise() |>
  mutate(rentals=nrow(titles)) |>
  mutate(unique_titles=list(distinct(titles))) |>
  select(-titles) |>
  arrange(customer) |>
  print(n=Inf, width=Inf)
```

The code for this query works with nested data, that is, data with columns that contain lists, created using the `nest()` function. In this example, `nest(titles=title)` creates a column called "titles" that contains a list of all the elements of the "title" column for each customer. The R code also demonstrates row-wise operations. Both `mutate()` functions after `rowwise()` function operate by row. Specifically, the first use of the `mutate()` function creates a new column "rentals" which contains the number of rows in the titles column *for each row* (recall that the "titles" column contains lists of film title). Similarly, the second use of the `mutate()` function creates a new column "unique_title" that contains a list of distinct film titles from the "titles" column *for each row*.

5.7 SQL and R

The "sqldf" library in R allows users to perform SQL queries on R data frames. Essentially, it provides a bridge between SQL and R. This integration allows users who are familiar with SQL to leverage its powerful querying capabilities directly on R data structures, without the need to switch between different tools or environments.

One of the main advantages of "sqldf" is its ability to handle large data frames more efficiently than some of R's native functions. By utilizing SQL queries, users can perform complex data manipulations and aggregations with ease. The package supports various SQL commands including SELECT, JOIN, ORDER BY, and GROUP BY, among others, enabling a wide range of data operations that are familiar to SQL users.

Under the hood, "sqldf" operates by temporarily converting data frames into databases, typically by creating an in-memory SQLite database, or, alternatively, using an existing database connection to any of a variety of RDBMS such as PostgreSQL. It then creates a table for each data frame, moves the data to the database tables, and executes SQL statements. It then moves the result set back into R as a data frame. This seamless process allows for a smooth integration of SQL's data processing capabilities within the R environment.

"sqldf" is particularly useful for R users who are already comfortable with SQL syn-

tax and for complex data manipulation tasks that might be more cumbersome or less intuitive in R's native syntax. Its ability to handle data frames as if they were SQL tables makes it a highly valuable tool for data analysts and statisticians who work with large datasets and require the flexibility and power of SQL within the R programming environment.

The following R code block shows a very simple example. Note that the SQL FROM clause recognizes data frame names; any columns used in the SQL query must be named columns from those data frames.

```
library(sqldf)
result_df <- sqldf('select distinct(title) from full_data')
```

When faced between the choice of data analytics using an SQL RDBMS or R/Tidyverse, there are a number of issues to consider:

- *Size of data*: R is limited by the amount of main memory of the computer. While large computers may offer 128GB or more, modern RDBMS can scale massively larger, in particular when distributing databases across a cluster of computers.
- *Access speed*: RDBMS have sophisticated indexing of tables and query planners that optimize complex queries for performance. While a dplyr analysis pipeline can also be optimized by carefully considering the order of function calls, the onus is on the data analyst to do this, while an RDBMS offer this "out-of-the-box".
- *Currency*: Using an RDBMS means that analytics can be performed on operational data, that is, the most current and up-to-date data. In contrast, the use of R involves first exporting data from the operational system and then analyzing it at a later time. However, while tempting, it is not generally recommended to perform complex analytics on an operational database, as it can significantly affect performance.
- *Transactions*: An RDBMS ensures consistent views of data across multi-user, concurrent updates. This means that, when using an operational database, the analysis sees consistent data, whereas an exported snapshot of the data may not necessarily be consistent, depending on the export mechanism.
- *Tools*: R has tools for statistical analysis and visualization, beyond mere reporting. So far, we have considered only simple descriptive analytics. However, when the data is to be used for sophisticated statistics or predictive analytics, it is no longer possible to do this on RDBMS.

These issues motivate the following recommendations:

- Do not "hit" operational RDBMS for heavy-weight or frequent analytics. While it may be fine to do the occasional summary analytics on an operational database, this should not be normally done.

- Regularly export consistent data from RDBMS. If up-to-date data is needed, automate the export from the database to occur at regular intervals. However, note also that exporting data has a performance impact on operational databases.
- Sometimes, SQL may be the more intuitive language to specify the required analysis. In these cases, use separate in-memory or on-disk RDBMS for analytics (e.g. with `sqldf`) if desired/required.
- Finally, if the size of data is too large to handle in R, consider distributed tools such as Hadoop/Spark that are made for Big Data analytics.

Hands-On Exercises

The following hands-on exercises are designed to familiarize you with the Tidyverse packages, especially the `dplyr` package. Use these exercises with the Pagila CSV data set.

1. Find all films with a rating of 'PG'
2. List all customers who live in Canada (with their address)
3. Find the average *actual* rental duration for all films
 - This requires date arithmetic, use the `lubridate` package
4. Find the average overdue time for each customer
 - This requires date arithmetic, use the `lubridate` package
5. List all films that have never been rented
6. List the names of actors who have played in more than 15 films

Chapter 6

Introduction to Data Management with Python

6.1 Introduction

Python is a high-level, interpreted programming language known for its simplicity and readability. It was created by Guido van Rossum and first released in 1991. Python's design philosophy emphasizes code readability through the use of significant¹ whitespace. This unique approach has contributed to Python becoming one of the most popular programming languages in the world.

Python's standard library of functions is large and comprehensive, covering a range of programming needs including web development, data analysis, artificial intelligence, scientific computing, and more. Its simplicity and versatility allow programmers to express concepts in fewer lines of code compared to languages like C++ or Java. Additionally, Python supports multiple programming paradigms, including procedural, object-oriented, and functional programming.

One of the biggest advantages of Python is its strong community support and the vast availability of third-party packages, which extend its capabilities even further. Frameworks like Django for web development, Pandas for data analysis, and TensorFlow for machine learning are just a few examples of Python's extensive ecosystem.

Python's popularity can be attributed to its wide range of applications in various fields, such as web development, data science, artificial intelligence, scientific computing, and scripting. It's often used in academic and research settings due to its ease of learning and its ability to handle complex calculations and data manipulation. Major tech companies and organizations use Python, showcasing its reliability and robustness.

¹"Significant" in this context does not mean lots, it means that spaces at the beginning of a line, that is, line indentations, have meaning and Python code does not work the same way without those spaces.

In terms of benefits, Python is known for its efficiency, reliability, and speed of development. It is often used for rapid prototyping and iterative development. Python's syntax is clean and its code is generally more readable and maintainable compared to many other programming languages. This readability makes it easier for developers to work on projects collaboratively.

Overall, Python's combination of versatility, simplicity, and powerful libraries makes it a preferred choice for both beginners and experienced developers across diverse fields. Its continued evolution and adaptation to new technologies and paradigms ensure its relevance in the fast-paced world of software development.

Intro Tutorial: A very good introduction to Python can be found at <https://python.swaroopch.com/>, or, as a downloadable PDF, at <https://github.com/swaroopch/byte-of-python/releases/>

6.2 Python versus R

Python and R are two of the most popular programming languages used in data science, each with its unique strengths and applications. Python, known for its general-purpose nature, offers a more comprehensive approach to business analytics, allowing not just data analysis and visualization, but also the integration of data science processes into web applications, production systems, and more. Its simplicity and readability make it a go-to language for a wide range of developers, including those who are not primarily data scientists.

Python's extensive libraries like Pandas for data manipulation, NumPy for numerical computations, Matplotlib and Seaborn for data visualization, and Scikit-learn for machine learning make it a powerful tool for business analytics. Moreover, Python's capabilities in machine learning and deep learning, with libraries like TensorFlow and PyTorch, make it a preferred choice for cutting-edge applications in AI.

On the other hand, R, originally designed for statistical analysis, is highly specialized in statistical modeling and data analysis. It offers a rich ecosystem of packages for statistical procedures, classical statistical tests, time-series analysis, and data visualization. R is particularly favored for its advanced statistical capabilities and its powerful graphics for creating well-detailed and high-quality plots.

The choice between Python and R often comes down to the specific requirements of the project and the background of the business analytics team. Python is generally more versatile and better suited for integrating business analytics into larger production applications. It is also the more popular choice for machine learning projects. R, meanwhile, is excellent for pure statistical analysis and visualizing complex data sets. It's often preferred in academia and research settings where complex statistical methods are more commonly required.

Both languages have strong community support and a wealth of resources, making

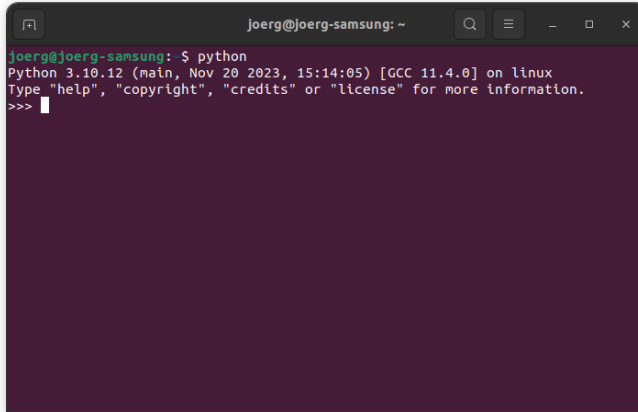


Figure 6.1: The Interactive Python Shell

them continually evolving tools in the field of business analytics. Many business analysts are proficient in both, choosing the one that best fits the task at hand. In collaborative settings, it's not uncommon to see teams utilizing both Python and R, leveraging the strengths of each to achieve more comprehensive and powerful data analysis outcomes.

6.3 Using Python

The Interactive Python Shell, Jupyter Notebooks, and PyCharm IDE represent different environments for Python development, each with distinct features and use cases.

Interactive Python Shell The Python shell is the most basic and straightforward environment for Python programming. Users can type Python code and see the results instantly. The simplicity is the primary advantage of the Python shell. The immediate feedback makes it excellent for experimentation, learning Python syntax, and quick tests. There is no need for creating files or setting up a project environment. This feature is especially beneficial for beginners who are just starting to learn Python, as it provides a straightforward way to test out new concepts and functions without the overhead of more complex development environments. Figure 6.1 shows a screenshot of the Python shell.

While the Python shell supports all the features of the Python language. However, it lacks advanced features found in full-fledged Integrated Development Environments (IDEs), such as code completion, debugging tools, or project management, which are essential for larger projects. Its simplicity is both a strength and a limitation: while it is easy to use, it might not be the best choice for larger programming projects.

On Ubuntu Linux, simply type `python` in a terminal window to launch the Python

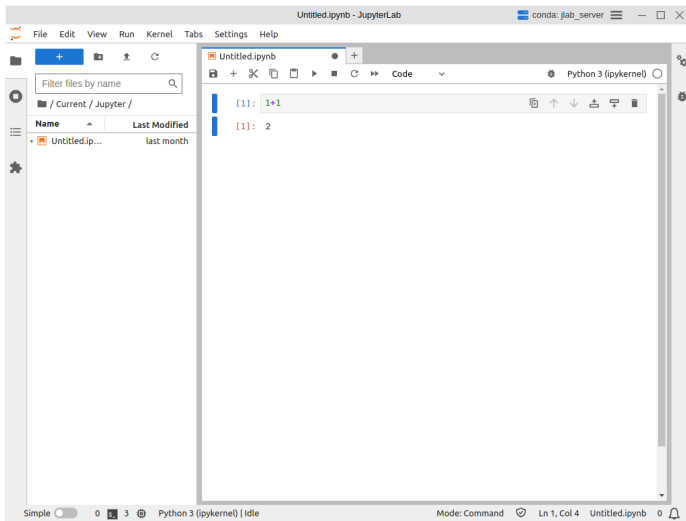


Figure 6.2: Jupyter Notebook

shell. On Windows and MacOS systems, you will find applications to launch the Python shell in a window. The shell prompts you for commands with the `> > >` prompt. Simply enter the command and press the `ENTER` key to execute a command. Use the `quit()` function to exit the shell. The Python shell remembers your previous commands, so you can use the up and down arrow keys to recall commands and edit them. The Python shell also performs code completion using the `TAB` key, which helps speed up coding and reduce typing errors. Similar to an R session, you should use a notepad editor application to assemble commands and then copy/paste them into the Python shell, as copy/paste results into a notepad editor. This makes editing long commands easier and ensures that your analysis will be repeatable.

Jupyter Notebooks Jupyter notebooks offer a more interactive and versatile platform, particularly favored in data science and academic research. Jupyter Notebooks allow users to create and share documents that can contain "cells" where each cell may contain Python code, text (using Markdown), equations (using LaTeX), or visualizations. This mix of Python code, documentation, description, and results makes it ideal for data exploration, visualization, and complex analyses where explaining the process is as important as the code itself, allowing for a narrative approach to coding. While Jupyter Notebooks support various programming languages, they are predominantly used with Python. Figure 6.2 shows a screenshot of a Jupyter notebook in the JupyterLabs Desktop environment.

The immediate feedback upon code execution helps in quick hypothesis testing and data manipulation. Furthermore, the integration of rich media alongside code makes Jupyter Notebooks an excellent tool for creating comprehensive documentation, tutorials, and educational materials.

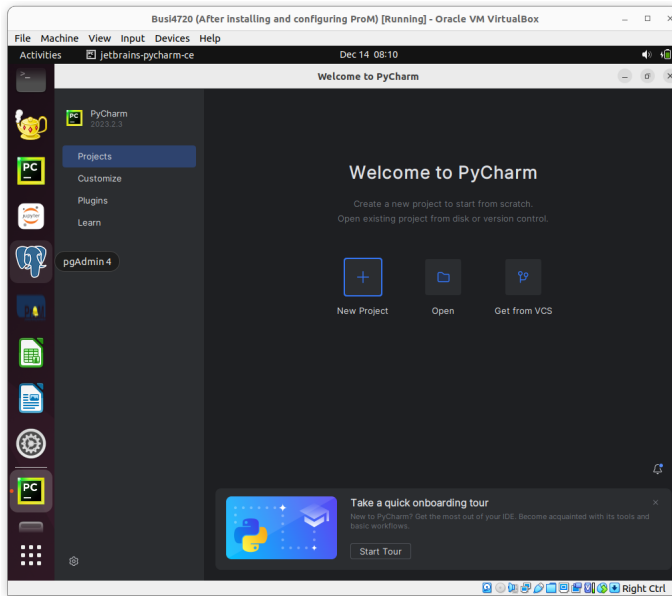


Figure 6.3: PyCharm Integrated Development Environment (IDE)

Notebooks can be easily shared, making them popular in collaborative projects. The ability to see the code, along with its output and accompanying explanation, in a single document enhances understanding and teamwork. Jupyter Notebooks run in a web browser, offering platform independence and eliminating the need for complex software application setups.

When working with Jupyter Notebooks, the term "kernel" denotes a particular version of the Python programming language and environment (i.e. Python packages, etc.) that runs your code. You can enter code in an empty cell and press **CTRL-ENTER** to execute code in the cell. A cell can contain multiple lines of code. Jupyter Notebook cells can be merged, split, moved, copied, and deleted, and you can save, import, and export notebooks, among much other, advanced functionality.

PyCharm IDE: The PyCharm Integrated Development Environment (IDE) is a full-featured software development environment designed specifically for Python. It offers a wide range of tools and features for professional software development, including code completion, debugging, project management, version control integration, and a powerful code editor. PyCharm is more suited for larger and more complex software projects. Its sophisticated environment, while powerful, might be overwhelming for beginners or for those who require a simple platform for exploratory data analysis. Figure 6.3 shows a screenshot of the PyCharm IDE.

One of the key strengths of PyCharm is its intelligent code editor, offering features like code completion, code inspections, and automated refactoring. These features greatly

enhance productivity and reduce the likelihood of programming errors. Additionally, PyCharm includes an integrated debugger and testing support, simplifying the process of diagnosing and fixing issues in programming code. The IDE also offers seamless integration with version control systems like Git, which is essential for collaborative development and code management.

In summary, while the Interactive Python Shell is best for quick, simple tasks and learning the basics, Jupyter Notebooks are ideal for business analytics projects that benefit from an interactive, explanatory, and exploratory approach. PyCharm is the most suitable for comprehensive software development, offering robust tools and features for managing complex codebases. The choice among these depends largely on the specific requirements of the project and the preferences of the developer.

6.4 Python Basics

The basic Python code in the following example prints a character string. The `print` function in Python is very versatile and provides different ways to print the values of multiple variables. In particular, character strings have a `format` function that can be used to substitute the {} placeholders with values, either by index/number, by name, or by position, as shown in the following Python code block that defines two variables, `age` and `name` and prints their values in a variety of ways:

```
print('hello world')

age = 19
name = 'Malina'
print('{0} is {1} years old'.format(name, age))
print('{name} is {age} years old'.format(name=name, age=age))
print('{} is {} years old'.format(name, age))
print(f'{name} is {age} years old')
print(name+' is '+str(age)+' years old')
```

Because Python commands can get quite long, Python allows for backslashes to break long lines and continue the command on the following line. While not needed in this case, the following code block illustrates how to use them:

```
print('This is a very long \
string and needs a second line')
i = \
5
print(i)
```

Multiline character strings are enclosed in triple quotes, and the line breaks form part of the string, as shown in the following example. Note that the `print(s)` function prints the line breaks in the character string.

```
s = '''This is line 1
and here is line 2
and now this is line 3'''
print(s)
```

As with R, you can use Python interactively as a calculator. It provides the usual arithmetic operators and comparison operators. The // operator is for integer division with floor (rounding down), the % operator is the modulus (remainder) operator. Boolean values are `True` and `False` in Python and *cannot* be abbreviated (unlike in R). The following Python code block illustrates typical usage:

```
2 + 2
2**4
13 // 3
-13 // 3
13 % 3
-25.5 % 2.25
3 < 5
3 > 5
3 == 5
(3 < 5) and (4 < 2)
(3 < 5) or not (4 < 2)
```

The next code example shows some useful string functions. The `startswith()` function does what its name suggests and returns a boolean (`True` or `False`) value. The `find()` function returns either the first position of a string in another string, or -1 if the string is not found.

```
language = 'Innuktitut'
if language.startswith('Innu'):
    print('Yes, the string starts with "Innu"')
if 'u' in language:
    print('Yes, it contains the string "u"')
if language.find('nuk') != -1:
    print('Yes, it contains the string "nuk"')
```

The `join()` and `split()` functions for character strings do as the their names suggest and work with Python lists, illustrated in the code block below:

```
# Joining and Splitting
delimiter = '_*_'
mylist = ['Nain', 'Hopedale', 'Makkovik', 'Rigolet']
mystring = delimiter.join(mylist)
print(mystring)
thelist = mystring.split(delimiter)
print(thelist)
```

Important: Note the use of leading whitespace or indentation in the lines after the `if` statement in the above code. In Python, this *whitespace is required for defining the program logic!* In the above example, the indented lines indicate the extent of the program block to be executed after the `if` statement. The normal leading whitespace is four spaces.

Lists in Python are ordered collections of items, and use square brackets [] as delimiters. Lists are mutable, i.e. their contents can be changed. Lists may contain items of different data types, including other lists or structured data types. Useful list functions are `len()` which returns the number of items in a list, `append()`, which adds items to the end of the list, and `sort()`, which sorts by value (only for compatible data types in the list). Items can be removed by position using the `del()` or by value using the `remove()` functions.

```
# Inuit deities
gods = ['Sedna', 'Nanook', 'Akna', 'Pinga']
print('There are', len(gods), 'deities:')
for god in gods:
    print(god, end=' ')
print()

# Appending to a list
gods.append('Amaguq')
print('\nThe list of deities is now', gods)

# Sorting a list
gods.sort()
print('The sorted list is', gods)

# Removing items from a list
print('The first deity is', gods[0])
olditem = gods[0]
del gods[0]
print('I removed', olditem)
print('The list is now', gods)
gods.remove('Pinga')
print('The list is now', gods)
```

The above example also shows the use of Python comments, beginning with # to the end of the line. The example also shows iteration ("repeating") with the `for` statement. Similar to the earlier example illustrating the `if` statement, note the required indentation (leading whitespace) in the line(s) after the `for` statement to indicate the extent of the code block that is repeated.

Tuples in Python are also ordered collections of items, but they are immutable, i.e. their contents cannot be changed. Tuples use round brackets () as delimiters.

```
# Inuit Nunangat
regions = ('Inuvialuit', 'Nunavut', 'Nunavik', 'Nunatsiavut')
print('Number of regions is', len(regions))

all_regions = 'NunatuKavummiut', 'Kalaallit', 'Inupiaq', regions
print('Number of all Inuit regions:', len(all_regions))
print('All Inuit regions are', all_regions)
print('Regions in Inuit Nunangat are', all_regions[3])
print('First region in Inuit Nunangat is', all_regions[3][1])
print('Number of all Inuit regions is', \
      len(all_regions)-1+len(all_regions[3]))
```

Important: Indexing in Python is zero based, that is, the first element in a list or tuple is number 0, while the last element is number `len()` - 1. This is in contrast to R, where indexing starts at 1.

Dictionaries (or short, "dicts") in Python are key-value pairs that map one element to another. In other programming languages, this data structure is also called a *map* or an *associative array* (because it associates keys with their values). Python uses curly brackets {} as delimiters; the keys and values are separated using :. The value for a key is retrieved using the square bracket operator []. Keys and values may be any data type.

```
# Largest cities
c = {
    'Inuvialuit': 'Inuvik',
    'Nunavut': 'Iqaluit',
    'Nunavik': 'Kuujjuaq',
    'Nunatsiavut': 'Nain'
}
print("Nunavik's largest city is", c['Nunavik'])
```

Keys and values can be retrieved separately using the function `keys()` and `values()`. Dicts are mutable, as the following example shows by removing an entry with `del` and adding another entry.

```

# Retrieving keys and values
print(list(c.keys()))
print(list(c.values()))

# Deleting a key-value pair
del c['Nunavut']
print('\nThere are {} cities left\n'.format(len(c)))
for region, city in c.items():
    print('{} is largest city of {}'.format(city, region))

# Adding a key-value pair
c['Nunavut'] = 'Iqaluit'
if 'Nunavut' in c:
    print("\nNunavut's largest city is", c['Nunavut'])

```

A useful function to create dicts from two lists is the `zip()` function, shown below. The `zip()` function creates an iterator over fixed-length tuples that are passed into the dictionary creation function `dict()` as key–value tuples:

```

towns = ['Hopedale', 'Makkovik', 'Nain', 'Postville', 'Rigolet']
pops = [596, 365, 1204, 188, 327]
pop_by_town = dict(zip(towns, pops))
print(pop_by_town)

```

In Python, lists, tuples, and character strings are examples of *sequences*. All sequences provide membership tests using `in` or `not in` operators, as shown in some of the examples above. Sequences also provide integer indexing and slicing. Note that the end index in a slicing expression is *not inclusive*, that is, the slice extends up to but does not include the final index. This makes it easy to write a slice like `[: len(a)]` where `a` is some sequence (rather than having to write `[: len(a) - 1]` as one would in R or other programming languages where the end index is inclusive).

The following code shows some examples for slicing tuples. Note the negative end index in the third example. A negative end index iterates from the end of a sequence forwards". The slice `regions[1:-1]` extends from the second element to the third of the four elements.

```

regions = ('Inuvialuit', 'Nunavut',
           'Nunavik', 'Nunatsiaavut')
# Slicing on a tuple
print('Item 1 to 3 is', regions[1:3])
print('Item 2 to end is', regions[2:])
print('Item 1 to -1 is', regions[1:-1])
print('Item start to end is', regions[:])

```

Slicing in Python is more advanced than slicing in R as not only the beginning and end index can be specified, but also the step size, as shown in the next Python code block.

The final example slices backwards.

```
# Slicing with step
print(regions[::-1])
print(regions[::-2])
print(regions[::-3])
print(regions[::-1])
```

Character strings are also sequences, and they support slicing or indexing the same way as other sequences in Python.

```
language = 'Innuktitut'
# Slicing on a string
print('characters 1 to 3 is', language[1:3])
print('characters 2 to end is', language[2:])
print('characters 1 to -1 is', language[1:-1])
print('characters start to end is', language[:])
```

In the above example, pay careful attention to the use of negative indices in the slicing expressions, both for the index as well as the step size.

Tip: To read and execute Python statements from a file, use the expression
`exec(open('filename.py').read())`

Hands-On Exercise

1. Create a *list* containing the numbers 1 to 10. Use list slicing to create a sublist with only the even numbers.
2. Using a *for* loop, sum all the items in the list.
3. Using a *for* loop, iterate over the list and print each number squared.
4. Write a program to append the square of each number in the range [1:5] to a new list.

Hands-On Exercise

1. Create a *tuple* with different data types (string, int, float).
2. Demonstrate how tuples are immutable by attempting to change its first element.
3. Write a program to convert the tuple into a list.

Hands-On Exercise

1. Create a *dictionary* with at least three key-value pairs, where the keys are strings and the values are numbers.
2. Write a Python script to add a new key-value pair to the dictionary and then print the updated dictionary.
3. Create a nested dictionary, that is, a dictionary whose values are dictionaries, and demonstrate accessing elements at various levels.

6.5 NumPy

NumPy, short for Numerical Python, is an essential package for the Python programming language, widely used for scientific computing and data analysis. It provides powerful numerical arrays and matrices, along with a large collection of high-level mathematical functions to operate on these arrays. The cornerstone of NumPy is its "ndarray" (n-dimensional array) object. These arrays are more efficient than Python's built-in lists, especially for numerical operations, due to their fixed type and contiguous memory allocation.

NumPy arrays facilitate advanced mathematical and statistical operations, including linear algebra, Fourier transform, and random number generation. The ndarray object supports vectorized operations, broadcasting, and indexing capabilities. This means that operations can be applied to entire arrays without the need for explicit loops, leading to cleaner and faster code.

One of the reasons for NumPy's popularity in the scientific and data science communities is its seamless integration with other Python libraries. Libraries like Pandas for data manipulation and analysis, Matplotlib for data visualization, and SciPy for scientific computing all build upon and work in conjunction with NumPy, creating a robust ecosystem for scientific computing tasks.

Tip: The NumPy website provides two very good introductions, in the form of the [Quick Start](#) and the [NumPy for absolute beginners](#) tutorials.

NumPy ndarrays have a set of useful properties or attributes, summarized in Table 6.1. Note that the terminology is "*axes*", rather than "*dimensions*" as in the previous chapter on R, although the `ndim` property of an ndarray uses the term "dimension" in its name.

The following Python code block illustrates the use of these properties. Note the use of the `arange()` function to create a one-dimensional array of 15 numbers (from 0 to 14), that is then reshaped into a 2-dimensional array with 3 rows and 5 columns. Rows are axis 0, and columns are axis 1.

<code>ndarray.ndim</code>	Number of axes
<code>ndarray.shape</code>	Type describing the size of each axis (dimension)
<code>ndarray.size</code>	Total number of elements
<code>ndarray.dtype</code>	The datatype of the elements, for example <code>numpy.int32</code> , <code>numpy.int16</code> , <code>numpy.float32</code> , or <code>numpy.float64</code>
<code>ndarray.itemsize</code>	Number of bytes for each element

Table 6.1: Attributes of NumPy ndarray

```
# Import the numpy package
import numpy as np

# Create an array
a = np.arange(15).reshape(3, 5)

print(a.shape)
print(a.ndim)
print(a.dtype.name)
print(a.size)
print(type(a))
```

The following Python code block shows element-wise operations and array operations on a NumPy array. Python determines automatically which functions are array functions (like `sum()`) and which ones are element-wise functions (like `sqrt()`). Note the creation of the array with the `array()` function from a list of two tuples. Note also the use of the `axis` parameter in the `max` function to specify whether to aggregate by row or by column. The `axis` parameter can also be applied to other functions like `sum()` or `std()`.

```
# Create an array from Python lists and tuples
b = np.array([(1.5, 2., 3), (4, 5, 6)])

# Elementwise operations
print(3 * b)
print(b + 5)
print(np.sqrt(b))

# Array operations
print(np.max(b))
print(np.max(b, axis=0))
print(np.max(b, axis=1))
print(np.std(b))
print(np.cov(b))
print(np.sum(b))
```

In the above example, the `std()` function without `axis` parameter computes the standard deviation of all elements in the array, while `cov` treats each *row* of the array as a vector and computes their variances and covariances. To treat array columns as vectors, either transpose the array first, using the `T` operator or use the `rowvar=False` parameter for the `cov()` function.

To create pre-initialized arrays, NumPy provides two convenience functions to create arrays filled with 0s or 1s:

```
# Create an array of zeros with shape (3, 4)
x = np.zeros((3, 4))
print(x)

# Create an array of ones with shape (2, 3, 4)
y = np.ones((2, 3, 4))
print(y)
```

In a generalization of the slicing expressions for Python sequences, each axis of a NumPy array can be sliced using the `[:] or [::]` expressions, as shown in the following example of a two-dimensional array. The slicing expressions for different dimensions are separated by commas.

```
b = np.array([[ 0,  1,  2,  3],
              [10, 11, 12, 13],
              [20, 21, 22, 23],
              [30, 31, 32, 33],
              [40, 41, 42, 43]])
print(b[2, 3])
print(b[0:5, 1])
print(b[:, 1])
print(b[1:3, :])
print(b[-1])
```

When not all axes are supposed to be sliced, one can omit initial or final unsliced axes in the slicing expression using the ellipsis "...” as shown in the following Python code block.

```
c = np.array([[[ 0,  1,  2],
               [ 10, 12, 13]],
              [[100, 101, 102],
               [110, 112, 113]]])
print(c[1, ...])
print(c[1, :, :, :])
print(c[..., 2])
print(c[:, :, :, 2])
print(c[..., :, :, 1])
```

NumPy arrays also provide convenient iteration of their rows and their elements. Note

the use of the `flat` operator to "flatten" a multi-dimensional array to a single dimension in the code block below.

```
for row in b:
    print(row)

for element in b.flat:
    print(element)
```

NumPy provides an easy way to reshape arrays to any dimension. However, it is important to be aware of where and how the elements move during such a reshape. The order can be specified using an optional argument to `reshape`; consult the NumPy documentation for details. The following example also demonstrates the use of the default random number generator² (`rng`) in NumPy to create an array of shape `(3, 4)` filled with random numbers between 0 and 1.

```
# Create a random number generator with seed 1
rg = np.random.default_rng(1)

# Create an array of shape (3, 4) of random numbers
a = np.floor(10 * rg.random((3, 4)))

# Show information about the array and reshape
print(a.shape)
print(a.flatten())
print(a.reshape(6, 2))
print(a.T)
print(a.T.shape)
```

The above example uses the `flatten()` function which returns a one-dimensional array, whereas the `flat` property returns an iterator to be used in a `for` loop. The `T` property returns the transpose of the array. In two dimensions, the transpose swaps rows and columns. The NumPy transpose is also defined for more than two dimensions, the axes are transposed such that `a.T.shape==a.shape[::-1]`.

The next example illustrates concatenation or stacking operations to stack two arrays either vertically, that is, by row, or horizontally, that is, by column. The arrays must be of compatible shape for these stacking operations.

```
b = np.floor(5 * rg.random((3, 4)))
print(np.vstack((a, b)))
print(np.hstack((b, a)))
```

²A random number generator in computer science is always a pseudo-random number generator that creates a sequence of numbers according to a deterministic formula (because computers are deterministic), starting from an initial "seed" number. The sequence is repeatable when beginning with the same seed. A good pseudo-random number generator will create sequences that are indistinguishable from true random numbers, for example, those created by rolling dice.

Arrays can be indexed also by boolean arrays. For example, in the following Python code block, the expression `a < 5` constructs a boolean array whose entries are `True` when the corresponding element in `a` is less than 5. This boolean array is then used to select or index the array `a`:

```
a = np.array([[1, 2, 3, 4],
              [5, 6, 7, 8],
              [9, 10, 11, 12]])
print(a[a < 5])
print(a < 5)
print(a[a%2 == 0])
print(a%2 == 0)
```

Finally, NumPy provides easy ways to identify unique elements in an array and to count how often particular elements occur in an array. The following example also demonstrates another use of the `zip()` function, already introduced above, to construct a list of tuples.

```
a = np.array([11, 11, 12, 13, 14, 15, 16,
              17, 12, 13, 11, 14, 18, 19, 20])
print(np.unique(a))

# Return the first index of a unique value
values, indices = np.unique(a, return_index=True)
print(list(zip(values, indices)))

# Return the counts of each unique value
values, counts = np.unique(a, return_counts=True)
print(list(zip(values, counts)))
```

Hands-On Exercises

1. Create an array with random numbers in the shape indicated by the last four digits of your student number (if your student number contains a 0, use a 1 instead)
2. Construct a new array by swapping the first half of rows (axis 0) with the second half of rows (axis 0)
3. Calculate all covariance matrices formed by the last two axes of your array. *Tip:* Iterate over the first two axes/dimensions with a `for` loop
4. Subtract the mean of the array from each element in the array (mean normalization)
5. Select all elements that are greater than the overall mean
6. Sort the selected elements from the previous step

6.6 Data management with Pandas

Pandas is a Python package widely used in data science, data analysis, and machine learning. It is known for its powerful data manipulation and analysis capabilities. It provides fast, flexible, and expressive data structures designed to make working with structured (tabular, multidimensional, potentially heterogeneous) and time series data both easy and intuitive.

Pandas is useful for data cleaning, data transformation, and data analysis. It offers functions for reading and writing data in various formats such as CSV, Excel, JSON, and SQL databases. The Pandas package simplifies handling missing data, merging and joining datasets, reshaping, pivoting, slicing, indexing, and subsetting data. Its time series functionality is particularly robust, offering capabilities for date range generation, frequency conversion, moving window statistics, date shifting, and lagging.

The library's design and functionality are heavily influenced by data analysis needs in finance, which is evident in its powerful group-by functionality for aggregating and transforming datasets, as well as its high-performance merging and joining of datasets. As part of the broader Python scientific computing ecosystem, which includes libraries like NumPy, Matplotlib, and Scikit-learn, Pandas plays an important role in data analysis and machine learning workflows.

Tip: The Pandas website provides very good [10 Minutes to Pandas](#) introductory tutorial for Pandas.

Central to Pandas are two primary data structures: the DataFrame and the Series. A Series in Pandas is a one-dimensional array-like object that can hold any data type, including integers, floats, strings, and Python objects. A DataFrame in Pandas is a two-dimensional, size-mutable, and potentially heterogeneous tabular data structure with labeled axes (rows and columns).

The following Python code constructs a Pandas Series of random numbers. The axis labels of a Series (and a DataFrame) are called "index" and allow one to name the elements. The example also shows how a Python dict can be converted into a series with named elements.

```
# Import the Pandas package
import pandas as pd

# Create a series from a NumPy array of random numbers
s = pd.Series(np.random.randn(5))
print(s.index)

# Provide indices (labels) when creating the series
s = pd.Series(np.random.randn(5), index=["a", "b", "c", "d", "e"])
print(s.index)

# Create a series from a Python dict that provides labels and values
d = {"a": 0.0, "b": 1.0, "c": 2.0}
print(pd.Series(d))

# Create a series from a dict and reorder the entries
print(pd.Series(d, index=["b", "c", "d", "a"]))
```

Note that in the last line of the above example, renaming or reordering the elements of the Series d introduces a NaN element for the index "d", because the dict contains no value for the key "d".

Pandas series behave largely like NumPy arrays, but note that to access their elements by numerical index, one has to use the `iloc` operator, as shown in the following Python code block. This allows slicing the same way as for Python sequences or NumPy arrays. The following example also shows that Series can behave like a Python dict, in that values for a named index ("key") can be retrieved. Series also provide membership tests for "keys" using `in`.

```
# Series behave like an ndarray
print(s.iloc[0])
print(s.iloc[:3])
print(s[s > s.median()])
print(s.iloc[[4, 3, 1]])
print(np.exp(s))

# Series behave like a dict
print(s['a'])
print(s['e'])
print('e' in s)
print('f' in s)

# Series have a datatype and name
s.name = 'My First Series'
print(s.dtype)
```

Pandas *DataFrames* are two-dimensional objects. Their columns may have different data types. Conceptually, DataFrames can be considered as a dict of Pandas Series, as the following example demonstrates.

```

d = {
    "one": pd.Series([1.0, 2.0, 3.0],
                     index=['a', 'b', 'c']),
    "two": pd.Series([1.0, 2.0, 3.0, 4.0],
                     index=['a', 'b', 'c', 'd'])
}
df = pd.DataFrame(d)
print(df)
print(df.index)
print(df.columns)
print(pd.DataFrame(d, index=['d', 'b', 'a'],
                    columns=['two', 'three']))

```

Constructing the DataFrame `df` "lines up" the two Series on their common indices, and will introduce a "NaN" for index "d" in column "one", because that Series does not contain a value for "d". Similarly, inserting a column named "three" in the last line of the above example yields a column filled with "NaN" because the dict `d` does not contain values for the key "three".

DataFrame columns can be accessed using their quoted name, and will yield a Pandas Series with the usual operations. The following Python code example shows that new columns can be added simply by defining them, as in the "flag" column below or using the `assign()` function, which works similarly to the `mutate` function in R/dplyr. Columns can be removed using the `del` command or the `pop()` function. The latter returns the deleted column as a Series.

```

print(df['one'])
df['three'] = df['one'] * df['two']
df['flag'] = df['one'] > 2
print(df)

del df['two']
three = df.pop('three')
df['foo'] = 'bar'
df['one_trunc'] = df['one'][:2]
df.insert(1, 'bar', df['one'])
print(df)

# Similar to 'mutate' in R/Dplyr
df = df.assign(four = df['one'] * np.sqrt(df['bar'])))
print(df)

```

Pandas DataFrames can be indexed by column, by label, by integer location, or by boolean vectors. Table 6.2 shows an overview of the different methods and their return values.

As noted earlier, Pandas automatically aligns data by indices, that is, by row and column labels, for operations on dataframes. Note how the addition of two dataframes of unequal shape introduces "NaNs". For convenience, NumPy operations can also be

Select column	<code>df['colname']</code>	Series
Select row by label	<code>df.loc['label']</code>	Series
Select row by integer location	<code>df.iloc[loc]</code>	Series
Slice rows	<code>df[::]</code>	DataFrame
Select rows by boolean vector	<code>df[bool]</code>	DataFrame

Table 6.2: Methods for indexing Pandas DataFrames

used to operate on Pandas DataFrames, which are automatically converted to NumPy ndarrays before and converted back after such an operation.

```
df = pd.DataFrame(np.random.randn(10, 4),
                  columns=["A", "B", "C", "D"])
df2 = pd.DataFrame(np.random.randn(7, 3),
                   columns=["A", "B", "C"])
print(df + df2)

# Elementwise operators
print(df * 5 + 2)
print(1/df)
print(df**4)

# Transpose
print(df.T)

# Using Numpy functions
print(np.exp(df))
print(np.asarray(df))
```

To apply element-wise character string operations on Series or DataFrames it is useful to use the `str` property:

```
# String functions with 'str'
s = pd.Series([
    "A", "B", "C", "Aaba", "Baca", np.nan,
    "CABA", "dog", "cat"], dtype="string")
s.str.lower()
```

Pandas provides a number of useful functions to get information about the contents of a DataFrame. The `info()` function provides information about the columns and their data types, while `head()` and `tail()` print the first and last few lines of a DataFrame.

```
df.info()
df.head()
df.tail(3)
```

The *boolean reduction* functions `all()` and `any()` operate by column on DataFrames with boolean values. As their names suggest, `all()` returns True when all entries in a column are true, whereas `any()` returns True if any of the entries in a column are true. The last line of the following Python code block re-applies `any()` to the Series that results from the first application of `any()`.

```
# Boolean reductions
(df > 0).all()
(df > 0).any()
(df > 0).any().any()
```

When making comparisons on DataFrames that include "NaN", it is important to realize that two "NaNs" are not equal when using the `==` operator, but they are equal when using the `equals` function. The following example illustrates this difference.

```
# NaN's are not the same
df.iloc[0,0] = np.nan
(df+df == df*2).all()
(df + df).equals(df*2)
```

Pandas provides useful functions for basic descriptive statistics and aggregation on DataFrames. In particular, the `describe()` function is useful to get a basic information on the data in a DataFrame. The `mean()` function takes as its optional first argument the axis number (0 for rows, 1 for columns) and can skip missing values when summing. Multiple aggregates can be formed using the `agg()` function. The Python code block below illustrates the use of these functions.

```
# Descriptive statistics
df.mean(0)
df.mean(1, skipna=False)
df_std = (df - df.mean()) / df.std()
df.describe()

# Aggregation with 'agg'
df.agg(['sum', 'mean', 'std'], 0)
```

Pandas DataFrames can be sorted by columns, and the functions `nlargest()` and `nsmallest()` can be used to select a DataFrame with only the n smallest or largest values in a given column.

```
# Sort by values
df.sort_values(by=['A', 'B'])
df.nsmallest(3, 'A')
df.nlargest(3, 'A')
```

A very useful way to identify or select data in a Pandas DataFrame is the `query()` function, which accepts a simplified boolean condition as argument. This allows one to write much shorter and compact selection logic, as shown in the following example. Note the two different forms of the same logical operator `&` and `and`.

```
df = pd.DataFrame(np.random.rand(10, 3),
                  columns=list('abc'))

# Pure python
df[(df['a'] < df['b']) & (df['b'] < df['c'])]

# Shorter with Query
df.query('(a < b) & (b < c)')
df.query('a < b & b < c')
df.query('a < b and b < c')
df.query('a < b < c')
```

The `query()` function can also be used for membership tests in Series and DataFrames using the `in` operator. This also is much more compact and easy to read than the pure Python `isin()` function. The following example code block shows the pure Python selection followed by equivalent selection with `query()`:

```
df = pd.DataFrame({'a': list('aabcccddeeff'),
                   'b': list('aaaabbccccc'),
                   'c': np.random.randint(5, size=12),
                   'd': np.random.randint(9, size=12)})

# Pure Python versus Query
df[df['a'].isin(df['b'])]
df.query('a in b')

df[~df['a'].isin(df['b'])]
df.query('a not in b')

df[(df['b'].isin(df['a'])) & (df['c'] < df['d'])]
df.query('a in b and c < d')

df[df['b'].isin(["a", "b", "c"])]
df.query('b == ["a", "b", "c"]')

df[df['c'].isin([1, 2])]
df.query('[1, 2] in c')
```

Pandas DataFrames also offer easy functions to remove duplicates. The following Python example code block shows how to identify rows that contain duplicate elements in a list of columns, and then remove the duplicates, keeping either the first or the last row. Note the different row indices in the retained results of `drop_duplicates` and their different values columns "c" and "d".

```
df2 = df.copy()

df2.duplicated(['a', 'b'])
df2.drop_duplicates(['a', 'b'], keep='last')
df2.drop_duplicates(['a', 'b'], keep='first')
```

Finally, Pandas provides many functions for reading and writing DataFrames from and to a variety of serialization formats and even SQL RDBMS. See the [Pandas IO user guide](#) for details.

6.7 The Pagila Database in Pandas

This section the use of Pandas for descriptive data analysis using the Pagila database data as an example. The Pagila database³ is a demonstration database originally developed for teaching and development of the MySQL RDBMS under the name Sakila⁴. Pagila is designed as a sample database to illustrate database concepts and is based on a fictional DVD rental store. It originally consists of several tables organized into categories like film and actor information, customer data, store inventory, and rental transactions. For this chapter, the Pagila data was summarized in a few related CSV files.

The following Python code block reads the rentals data of the Pagila database into a Pandas DataFrame using the `read_csv()` function. It then converts the data type of some columns from character strings to datetime types so that one can use date and time operations and arithmetic later.

```
# Read CSV
rentals = pd.read_csv('rentals.csv')

# Convert data types
rentals['rental_date'] = \
    pd.to_datetime(rentals['rental_date'], utc=True)
rentals['return_date'] = \
    pd.to_datetime(rentals['return_date'], utc=True)
rentals['payment_date'] = \
    pd.to_datetime(rentals['payment_date'], utc=True)

# Basic information
rentals.info()
rentals.describe()
rentals.index
rentals.columns
rentals.shape
```

³<https://github.com/devrimgunduz/pagila>,
<https://github.com/devrimgunduz/pagila/blob/master/LICENSE.txt>

⁴<https://dev.mysql.com/doc/sakila/en/>,
<https://dev.mysql.com/doc/sakila/en/sakila-license.html>

When working with data, it is often useful to first identify and remove missing values. The following Python code block first identifies columns (`axis=1`) in the dataset that contain any (`any()`) missing values (`isna()`). Of these filtered rentals, only some columns are selected.

```
filtered_rentals = rentals[rentals.isna().any(axis=1)]  
selected_rentals = \  
    filtered_rentals[  
        ['last_name', 'rental_date', 'return_date', 'title', 'amount']]
```

When printing DataFrames, Pandas by default abbreviates the output to manageable size. The number of rows and number of columns to be printed is controlled by two Pandas options that can be set as shown in the following example, which removes any limits.

```
pd.set_option('display.max_rows', None)  
pd.set_option('display.width', None)
```

The remainder of this section shows how Pandas can be used to provide equivalent results as obtained in the previous chapter using R/dplyr and in the chapter on relational databases with SQL. *Compare the Python code to the R code and the SQL code to achieve similar results.*

Example: Find all films and the actors that appeared in them, ordered by film category and year, for those films that are rated PG.

```

actors = pd.read_csv('actors.categories.csv')

result = pd.merge(rentals, actors, on='title',
                  suffixes=('_customer', '_actor'), how='outer')

result = result[result['rating'] == 'PG']
result['actor'] = result['last_name_actor'] + \
                  ', ' + result['first_name_actor']

result.rename(columns={'release_year': 'year'}, inplace=True)

result = result[['actor', 'title', 'category', 'year']]

result.drop_duplicates(['actor', 'title', 'category', 'year'],
                      inplace=True)

result.sort_values(['category', 'year', 'title'], inplace=True)

grouped = result.groupby(['category', 'year', 'title'])

g_result = grouped['actor'].apply(list).reset_index()

print(g_result)

```

This Python code block above performs a series of data manipulation operations using Pandas. The operations merge, filter, transform, and group data from the Pagila movie rental dataset.

- *Reading Data:* The code reads a CSV file named 'actors.categories.csv' into a DataFrame called "actors".
- *Merging DataFrames:* It then merges two DataFrames: "rentals" and "actors", based on the "title" column that is common to both DataFrames. The `suffixes` parameter is used to differentiate columns with the same name in both DataFrames, by adding either "_customer" or "_actor" to the column names. The `how='outer'` parameter ensures that all records from both DataFrames are included in the result, even if there are no matching titles in one of them.
- *Filtering Data:* After merging, the script filters the resulting DataFrame to include only rows where the "rating" column contains the value "PG".
- *Creating a New Column:* A new column, "actor", is created by concatenating the "last_name_actor" and "first_name_actor" columns, separated by a comma.
- *Renaming a Column:* The "release_year" column is renamed to "year".
- *Selecting and Rearranging Columns:* The DataFrame is then reduced and rearranged to include only the columns "actor", "title", "category", and "year".
- *Dropping Duplicates:* Duplicate rows based on the combination of "actor", "title", "category", and "year" are removed. This ensures that each combination is unique in the dataset.

- *Sorting Data:* The DataFrame is sorted by "category", then "year", and finally "title".
- *Grouping Data and Creating a List:* The data is grouped by "category", "year", and "title". For each group, the "actor" values are aggregated into a list. This creates a list of actors for each movie title, categorized by year and category.
- *Resetting Index:* After the grouping and aggregation, the index is reset to turn the grouped data back into a regular DataFrame.
- *Printing the Final Result:* Finally, the processed DataFrame is printed.

Example: Find the most popular actors in the rentals in each city.

The Python code block below combines the data frames from the multiple CSV files that make up the Pagila data set, because the combined, full data is used for other analysis examples below.

- The Python code merges the "rentals" DataFrame with the "addresses" DataFrame based on the columns "customer_address" in "rentals" and "address_id" in "addresses" to linking rentals with corresponding customer addresses.
- The script then merges the resulting DataFrame with the "actors", based on the "title" column.

```
addresses = pd.read_csv('addresses.csv')
addresses['phone'] = addresses['phone'].astype(str)

full_data = pd.merge(rentals, addresses,
                     left_on='customer_address',
                     right_on='address_id')

full_data = pd.merge(full_data, actors, on='title',
                     suffixes=('_customer', '_actor'))
```

The following Python code block performs the required analysis to on the full data constructed above, using the following steps:

- The code groups the data by "city" and "actor" and calculates the size of each group. This results in a count of how many times each actor's movies were rented in each city. The result is reset into a DataFrame "grouped" with a new column "count" representing these sizes.
- Within each city, actors are ranked based on the "count" column, with the ranking stored in a new column "ranking". The `rank` method is set to '`min`', which means actors with the same count will have the same rank, and it ranks in descending order of count.
- The code filters the DataFrame to select the top 3 actors (or ties) in terms of rental counts in each city.

- The filtered data is then sorted by "city", "ranking", and "actor" before being printed.

```
full_data['actor'] = full_data['last_name_actor'] + ', ' + \
    full_data['first_name_actor']

grouped = full_data.groupby(['city', 'actor']).size() \
    .reset_index(name='count')

grouped['ranking'] = grouped.groupby('city')['count'] \
    .rank(method='min', ascending=False)

filtered = grouped[grouped['ranking'] < 4]

sorted_df = filtered.sort_values(by=['city', 'ranking', 'actor'])

print(sorted_df)
```

Example: Find the customers who spend the most on rentals, and the number of rentals with the highest total rental payments for each category grouped by rental duration.

```
full_data['customer'] = full_data['first_name_customer'] + ' ' + \
    full_data['last_name_customer']

selected_data = full_data[['customer', 'amount', 'rental_duration', \
    'category', 'phone', 'city']]

grouped_data = selected_data \
    .groupby(['category', 'rental_duration', 'customer']) \
    .agg(amount=('amount', 'sum'), num_rentals=('amount', 'count')) \
    .reset_index()

grouped_data['ranking'] = grouped_data \
    .groupby(['category', 'rental_duration'])['payments'] \
    .rank(method='min', ascending=False)

top_entries = grouped_data.loc[
    grouped_data.groupby(['category', 'rental_duration'])['ranking'] \
        .idxmin()]

print(top_entries)
```

By now, it should be clear what most of the functions in the analysis accomplish. However, two important new things to note. First, the `agg()` function computes aggregates of the values in its first argument using the function in its second argument and stores the aggregate values in a new column. For example, the code below creates a new column "payments" with the "sum" of the values of the "amount" column of the grouped data, and a new column "num_rentals" with the "count" of the values of the "amount"

column. Second, the `idxmin()` function within the `loc[]` operator of the dataframe selects the smallest index, i.e. the smallest (highest) ranking when the data is grouped by category and rental duration.

Example: Get the top 5 and the bottom 5 grossing customers for each quarter.

This example demonstrates the Pandas date and time function `to_period()`. The argument `Q` returns quarters. Other frequently used arguments are `'D'` for days, `'W'` for weeks, `'M'` for months, `'Y'` for years, `'H'` for hours, and `'T'` for minutes. The use of the `sort_values()` function demonstrates "mixed" sorting, ascending by quarter, and descending by payments.

```
full_data['customer'] = full_data['first_name_customer'] + ' ' + \
    full_data['last_name_customer']

full_data['q'] = pd.to_datetime(full_data['rental_date']).dt \
    .to_period("Q")

selected_data = full_data[['customer', 'q', 'amount', 'rental_date']]

grouped_data = selected_data.groupby(['q', 'customer']) \
    .agg(amount=('amount', 'sum')).reset_index()

distinct_data = grouped_data \
    .drop_duplicates(['customer', 'q', 'payments'])

distinct_data['rank_top'] = distinct_data \
    .groupby('q')['payments'].rank(method='min', ascending=False)

distinct_data['rank_bot'] = distinct_data \
    .groupby('q')['payments'].rank(method='min', ascending=True)

filtered_data = distinct_data[(distinct_data['rank_top'] < 6) | \
    (distinct_data['rank_bot'] < 6)]

sorted_data = filtered_data \
    .sort_values(by=['q', 'payments'], ascending=[True, False])

print(sorted_data)
```

Example: Find the set of film titles by rental customer and the total number rentals for each customer.

The code below introduces *Lambda* functions. Lambda functions are unnamed, in-line functions, here it is a function that converts its parameter `x` to a set (i.e. it removes duplicates), and then converts the set to a list. The Lambda function is used as an argument to the `apply` function, that is, it is applied to all elements of the "titles" column in the grouped data. Recall that the "titles" column was introduced earlier in the script when the `list` function was applied to the "title" column of the grouped data and contains a list of film titles.

```
full_data['customer'] = \
    full_data['first_name_customer'] + ' ' + \
    full_data['last_name_customer']

selected_data = full_data[['customer', 'title']]

grouped_data = selected_data \
    .groupby('customer')['title'] \
    .apply(list) \
    .reset_index(name='titles')

grouped_data['rentals'] = grouped_data['titles'].apply(len)

grouped_data['unique_titles'] = grouped_data['titles'] \
    .apply(lambda x: list(set(x)))

grouped_data = grouped_data.drop(columns=['titles'])
sorted_data = grouped_data.sort_values(by='customer')

print(sorted_data)
```

Hands-On Exercise

1. Find all films with a rating of 'PG'
2. List all customers who live in Canada (with their address)
3. Find the average *actual* rental duration for all films
 - This requires date arithmetic
4. Find the average overdue time for each customer
 - This requires date arithmetic
5. List all films that have never been rented
6. List the names of actors who have played in more than 15 films

Chapter 7

Data Visualization in R and Python

7.1 Introduction

Data visualization, the practice of transforming information into a visual context, has become an indispensable part of modern data analysis and communication. This field intersects art and science, requiring both creativity and analytical skills to convert complex data sets into comprehensible, insightful visual representations. The motivations for visualizing data are multifaceted. Primarily, it enhances understanding by simplifying complex information, making patterns, trends, and correlations more apparent than they would be in raw data. It also aids in storytelling, where data-driven narratives can be compellingly presented to a broad audience, regardless of their expertise in data analysis.

The purpose of visualization is to simplify, summarize and abstract complex information into an easy to understand format for human consumption, for understanding, persuasion or explanation, or for decision making. Visualizations can help to compare different objects or things, they can help identify trends, patterns, and relationships. In general, visualizations help in understanding data and gaining insight into a domain or phenomenon.

The recent history of data visualization is marked by rapid advancements fueled by technology. In the last few decades, the advent of powerful computing and sophisticated software tools has revolutionized this field. Where once it was the domain of experts and specialists, data visualization has become accessible to a broader audience. Tools ranging from simple spreadsheet applications to advanced data visualization software have democratized the creation and interpretation of visual data. The rise of big data and machine learning has further escalated the importance of data visualization. As data sets have grown in size and complexity, the need for effective visualization

tools has become more pronounced, leading to innovative methods and approaches. Interactive visualizations, real-time data mapping, and the use of virtual and augmented reality are some of the cutting-edge trends redefining how we see and interact with data today. This evolution continues as we find new ways to visually interpret the vast and growing ocean of data that characterizes the digital age.

Visualization is important because humans are very good at visual pattern recognition. In fact, humans are too good at this, as they tend to also recognize patterns where none exist. This makes it easy to deceive oneself or others with data visualizations. Hence, visualization should always be undertaken with and supported by statistical data analysis.

Visual Discovery

Visual discovery refers to the use of interactive visualization tools to uncover hidden patterns, trends, and insights in data. This approach is a crucial aspect of modern data analysis, emphasizing the power of human visual perception. Visual discovery leverages the human brain's innate ability to process visual information rapidly. By translating complex data sets into graphical representations, it enables quicker and more intuitive understanding. Users can spot trends, outliers, and patterns more easily than they could through rows of numbers or text.

Visual discovery is a highly iterative and dynamic process. Analysts rapidly create or change data visualizations, such as charts, graphs, and maps, to explore different aspects of the data. This interactivity allows for real-time exploration and analysis, making it easier to drill down into specifics or zoom out for a broader view.

Visual discovery may be purely exploratory, without any prior knowledge by the data analyst, or it may seek to confirm or verify the beliefs or hypotheses that the data analyst has formed about the particular domain. However, even this confirmation is never final, but only a way to new insights and exploration. In this process, the analyst explores the data, forms some beliefs or hypotheses based on the exploration, tries to support it with a different visualization, and updates their beliefs or hypotheses based on the later visualization.

Declarative Visualization ("Storytelling")

In contrast to visual discovery, declarative visualization is purpose-driven and aims to provide explanations to a particular audience. It is *not* interactive or dynamic. Visualizations are intended to affirm or support a conclusion and to convince an audience or group of stakeholders. Information is not so much explored, as it is merely presented and explained in visualization. Declarative visualization is used to support decision making and is mainly static.

Operational Visualization (Monitoring)

In operational visualization, graphs and charts are used for supervision or monitoring of the operation of a system. They provide system supervisors or controllers with a real-time view of the state of key system properties and are used to spot situations or trends that require intervention in the system's operation, that is, operational decision making.

Quantitative Messages

Good visualizations are focused on the quantitative message they are intended to convey. For example, to present information about a time-series, that is, time-dependent behaviour of one or more variables, a line chart is a good type of visualization. However, that line chart would not be as useful to convey relative rankings of items or objects. For this purpose, a bar chart may be better suited. On the other hand, to describe part-whole relationships, a pie chart may be useful to show what part of the whole is contributed by its parts. Deviations from a mean or other standard, whether positive or negative, can be easily understood from a bar chart as well. To understand frequency distributions, one might use boxplots or histograms. Boxplots show median values, and measures of the "spread" or variability of the data. Histograms can show a one or two dimensional frequency distribution of values. To understand correlations of variables, a scatterplot is useful, where individual data points are plotted in a two or three dimensional coordinate system, often augmented with statistical information about their relationship. Finally, geographic information may use map data for visualization. This is sometimes called a "cartogram". Points may be overlaid on a map, or areas of a map may be colored or otherwise highlighted. In summary, it is important to consider the message to convey or the insight to be gained from a visualization when selecting the type of graph or chart.

7.2 Honesty in Visualization

For a number of reasons, it is easy to deceive with misleading visualizations. Humans are prone to see trends or patterns where none exist. A misleading visualization can use exploit this propensity in order to suggest relationships between objects or variables that do not exist. Humans recognize some aspects of visualization better, earlier, and easier than others. For example, humans recognize the length of a line easier than the area of a surface, and recognize variations in color better than they interpret textual labels. A misleading visualization can exploit these cognitive effects to make the interpreter focus on particular, misleading aspects in the visualization. Finally, because visualizations are intended to abstract from the data itself and provide a summary, visualizations may not include sufficient information about the data or its processing to allow the reader to understand what is shown, making it easy to suggest interpretations that are misleading.

Here are some general guidelines for using visualizations:

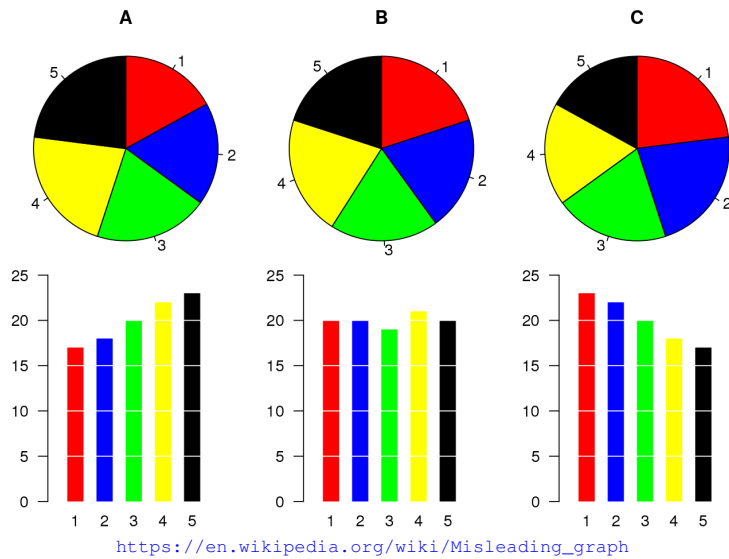
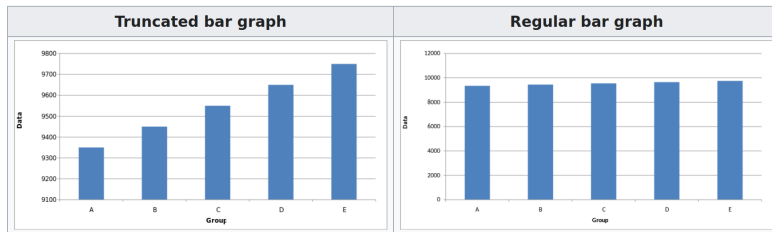


Figure 7.1: Comparing Pie Charts

- Do not deceive the target audience
- Do not diminish or hide relationships or trends
- Do not exaggerate relationships or trends
- Do not obfuscate, confuse, or hide information

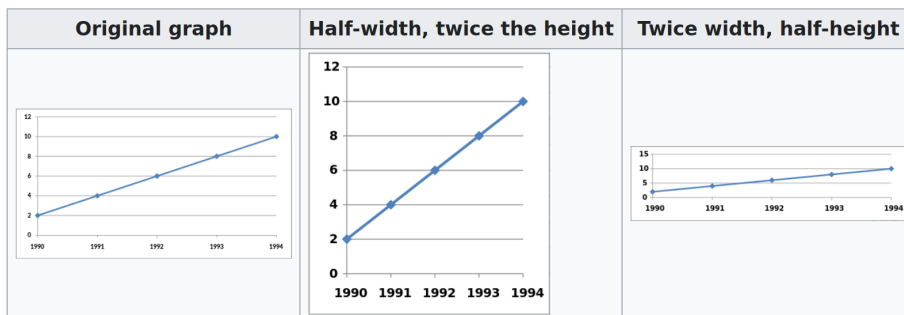
The term "dark pattern" has been coined to describe the opposite of best practices in a field, practices intended to deceive, mislead, or frustrate others. There are many of such dark patterns in visualizations:

- Use an inappropriate graph or chart type to hide or obfuscate relationships or trends. As noted above, different types of graph are suitable to convey different types of messages. An example is shown in Figure 7.1 that illustrates that a bar or column chart is more useful for comparisons of objects than pie charts, so the use of a pie chart could hide or obfuscate trends that may otherwise be prominent.
- *Graph unrelated data to suggest non-existent relationships.* The viewer of a visualization expects that data that is graphed together has a meaningful relationship. Simply by graphing data together, the analyst suggests a relationship where none may exist.
- *Scale multiple vertical axes to suggest correlations.* Scaling a graph with multiple vertical axes so that lines better align shows visual similarities that are not borne out by the data.
- *Use confusing colors.* For example, a color palette whose perceived color differ-



https://en.wikipedia.org/wiki/Misleading_graph

Figure 7.2: Truncated Axes



https://en.wikipedia.org/wiki/Misleading_graph

Figure 7.3: Scaling Axes and Aspect Ratios

ences do not map linearly to the actual differences in the data may be misleading. For another example, using different shades of the same color for values that are very different will visually diminish the difference.

- *Omit summary statistics.* For example, showing only the mean or median values, e.g. in a line or bar chart, omits the uncertainty in the data. It is better to also include error bars, information about quartiles or outliers in the chart to show variability or uncertainty, especially when there is significant uncertainty about differences or absolute values.
- *Truncate or scale axes to hide or exaggerate trend.* Truncating or scaling axes leads to increased slopes of lines or perceived differences between points or levels. This exaggerates differences or trends. Figure 7.2 shows an example of how small differences (right) can be exaggerated (left) in a bar chart. Similarly, Figure 7.3 shows how scaling or the use of different aspect ratios can be used to visually exaggerate or diminish trends or relationships between variables.
- *Scale in multiple dimensions.* The relative change or difference should be represented by a single dimension only. For example, in a bar or column chart, only the height or length of bar/column should change, not its width or area as well. This issue is often connected to the use of 3-dimensional graphics. While visually appealing, they exaggerate the apparent visual area of a foreground object,

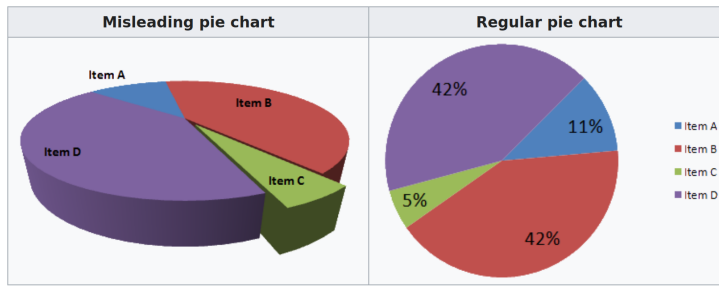


Figure 7.4: 3D Pie Charts

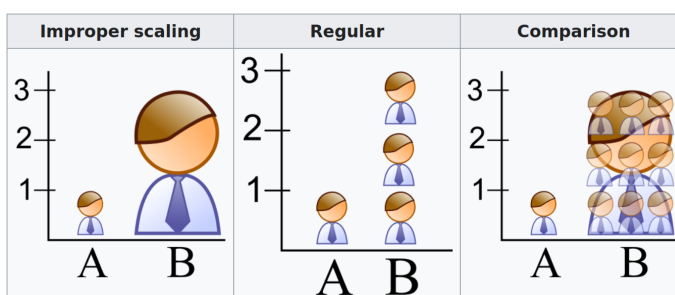
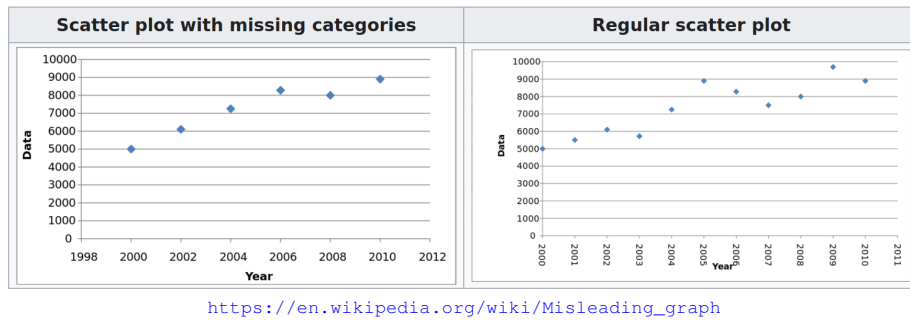


Figure 7.5: Scaling Multiple Dimensions

as illustrated in Figure 7.4. A related issue is the use of images in graphs, shown in Figure 7.5. In the improper scaling, the image is enlarged in two dimensions, suggesting a larger difference than there actually exists in the data.

- *Plot cumulative growth to hide trend.* A cumulative trend will always a positive trend, even as the contribution of individual items decreases sharply.
- *Use maps for non-geographic data.* Maps represent geographical area, rather than population or some other variables of interest. For example, coloring a map by voter preference visually overemphasizes thinly populated but large geographic areas.
- *Use incomplete data ("cherry-picking").* This includes examples such as showing only the previous year's data, instead of data for the previous five years to hide a trend, showing quarterly data instead of weekly data to hide volatility, or showing every data for every second month instead of for every month to hide specific data points or trends. Figure 7.6 shows an example of this dark pattern.
- *Use invalid data.* When data is known to be unreliable, that is, its quality is low, its uncertainty may be high, and it has a large error rate, it is misleading to use it to convey a quantitative message.



https://en.wikipedia.org/wiki/Misleading_graph

Figure 7.6: Incomplete Data

The comics in Figure 7.7, taken from the popular XKCD website¹ shows some of these visualization dark patterns in a humorous way.

In summary, misleading charts and visualizations can be particularly problematic because they exploit the visual nature of human perception, making the deception less noticeable. It is crucial for both creators and consumers of data visualizations to be aware of these pitfalls and to approach data representation and interpretation with a critical eye.

7.3 Special Types of Data and Visual Analytics

Streaming Data

Visualizing streaming data, also known as real-time data visualization, involves the dynamic representation of data that is continuously updated as new data arrives. This type of visualization is essential in contexts where timely and rapid data interpretation is critical, such as in financial trading, or network monitoring.

Streaming data presents some specific challenges for visualization. One of the primary challenges is managing the high velocity and volume of streaming data. The system must process and visualize data quickly enough to keep up with the incoming stream. Typically, only a limited window of data is available while older data is discarded. This means that, since the focus is on real-time data, it can be challenging to provide sufficient historical context for users to understand the current data in a broader temporal perspective. Moreover, due to the highly dynamic nature of the data, presenting streaming data in a way that is not overwhelming to the user is challenging. The visualization must strike a balance between providing enough detail and overloading the user with information, in particular information about changes, and between providing responsive graphs and overloading the user with such rapid changes they lose the ability to understand the data.

¹All XKCD comics are copyright by their creator (www.xkcd.com) and licensed under CC-BY-NC.

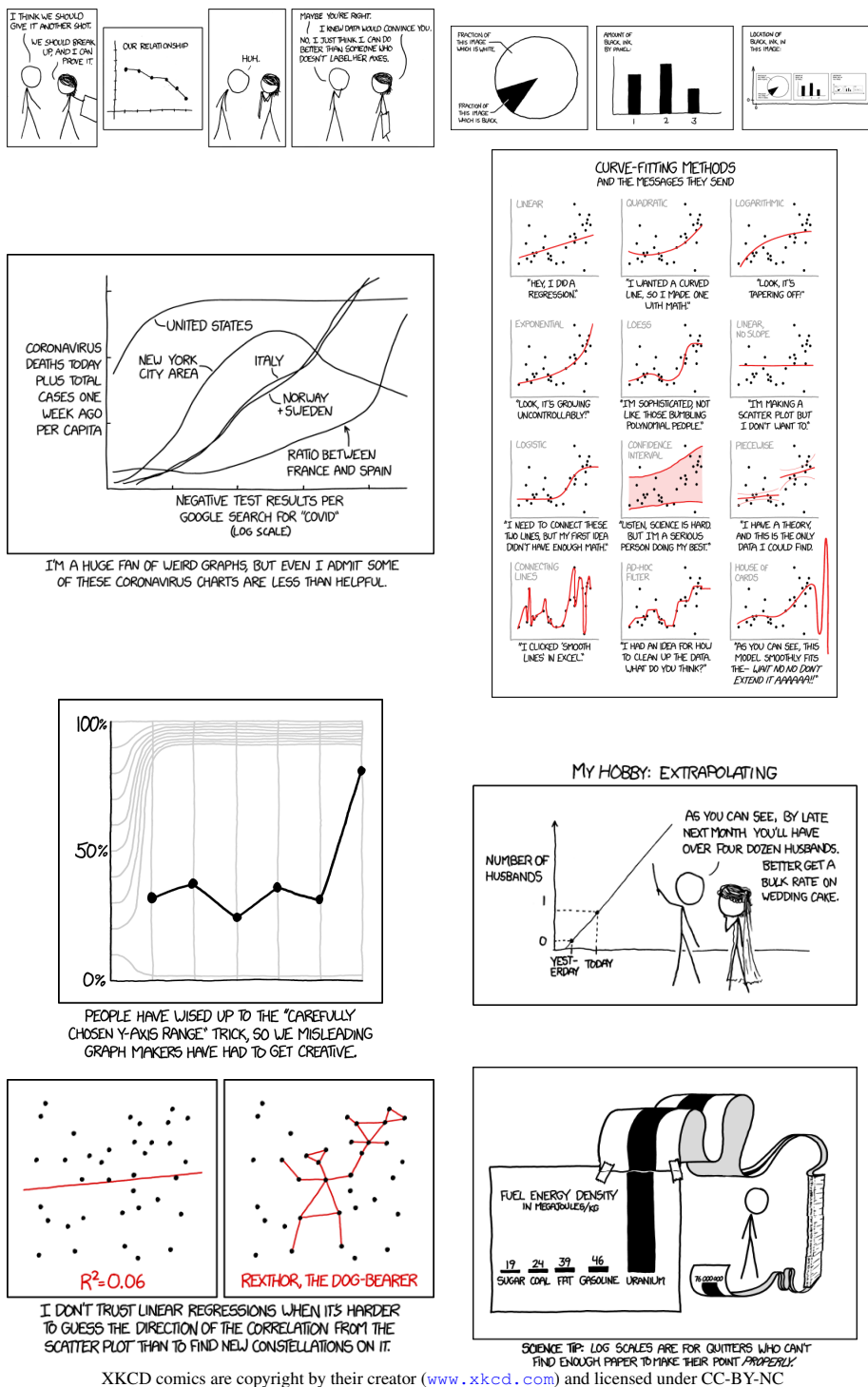
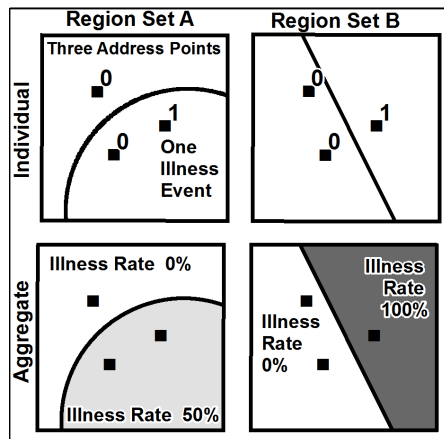


Figure 7.7: Visualization Comics by XKCD



https://en.wikipedia.org/wiki/File:Maup_rate_numbers.png

Figure 7.8: Different types of spatial divisions lead to different interpretations

Spatial Data

Visualizing geospatial or geographical data involves representing information that has a spatial component on a map or in a spatial context. While this type of visualization can be powerful for revealing patterns and insights related to location and geography, it presents some unique challenges. Geospatial data is often complex and multidimensional, encompassing not only locations but also attributes like time, elevation, population density, and more. Geospatial datasets can be very large, especially with the advent of satellite imagery, IoT (Internet of Things) sensors, and other sources of big data. Moreover, the granularity of physical space can range from very small areas of a few square meters to very large areas, such as provinces or states. For example, postal-code level data can produce very large data sets, even in small jurisdictions.

A specific problem is the choice of areal unit to use for data analysis or visualization. For example, location data points can be aggregated by counties or districts, by postal code areas, by school districts or school intake areas, by police or fire service coverage, or many others. Each of these different areal units will lead to different data summaries and therefore also to different visualizations. Choosing the type of area to use as the basis for visualization can have a large impact on the insights gained or the messages conveyed to the audience. A simple example is shown in Figure 7.8 that shows how aggregate statistics depend on the type of areal unit or boundary.

Another particular challenge with spatial data is mapping the three-dimensional Earth onto a two-dimensional surface. This mapping inevitably involves some form of projection, which can distort spatial relationships. Choosing an appropriate map projection that minimizes distortion for the specific data and use case is a critical challenge. There are many such projections², that distort or leave undistorted various properties such as

²https://en.wikipedia.org/wiki/Map_projection

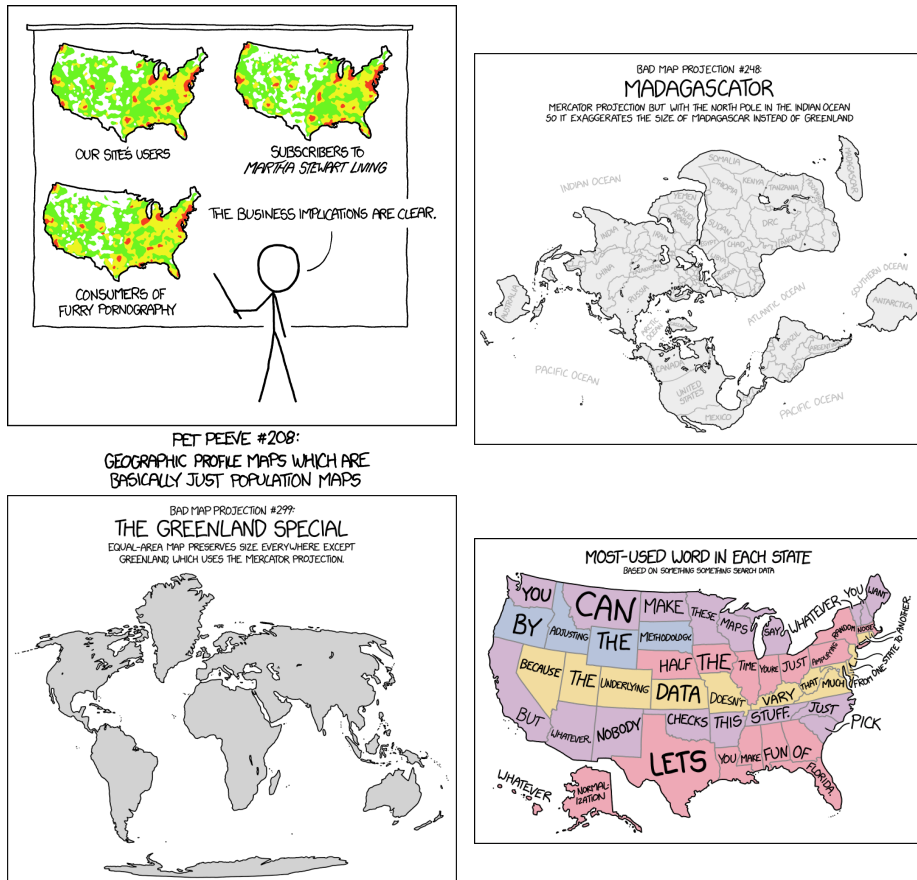


Figure 7.9: Map Visualization Comics by XKCD

lengths, areas, or angles. Figure 7.9 shows some of these issues in a humorous way.

Network and Graph Data

Visualizing network or graph data involves representing entities as nodes and the relationships between them as edges in a graphical format. This type of visualization is crucial for understanding complex structures in various fields like social network analysis, biology, computer science, and more. Typically, nodes are represented as boxes, circles, or textual labels, while edges are represented as lines or curves. Directed graphs use arrowheads on lines or curves to indicate the directionality of an edge.

Network visualization poses several unique challenges. Networks, especially large ones, can become very complex and cluttered when visualized. As the number of nodes and edges increases, the visualization can quickly become a tangled mess, making it

difficult to discern meaningful patterns or relationships.

To effectively explore graph data, especially large graphs, interactive features like zooming, panning, and highlighting are essential. Finally, graphs often contain large sets of attributes for nodes and edges. Representing these attributes effectively without cluttering the visualization or overwhelming the viewer is challenging. Techniques like color coding, sizing, or shaping nodes and edges are commonly used but require careful design.

In densely connected networks, edges can overlap, and nodes can occlude each other, leading to a loss of information and making it difficult to trace relationships or identify individual elements. Choosing an appropriate layout algorithm is therefore crucial for network visualization. There exist many different ways to visually layout a graph to make it visually clear and easy to understand and generally of high quality.

One of the most commonly-used types of algorithms position graph vertices based on physical metaphors of attractive and repulsive forces, for example an imaginary system of physical springs. Adjacent vertices are modelled with an attractive force, while all vertices have a repulsive force. The graph layout algorithm then tries to produce a layout in which an overall energy function is minimized. Figure 7.10 shows an example of such a graph layout.

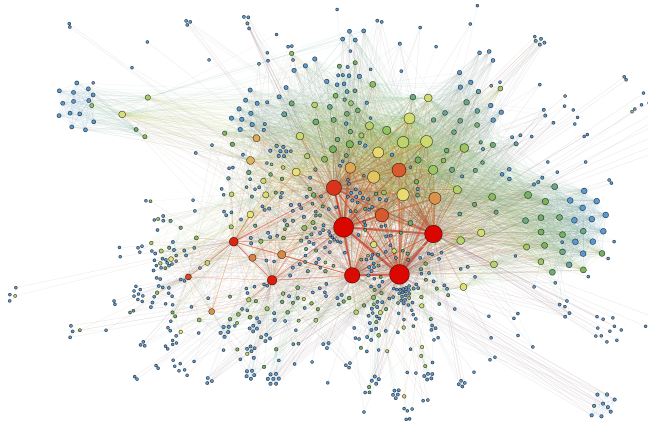
Another commonly used graph layout algorithm is the simple circular layout, where nodes are arranged equidistantly around a circle with edges drawn as lines or arrows. Figure 7.11 shows an example this type of graph layout.

In an arc diagram, as shown in Figure 7.12, the nodes are arranged on a straight line while edges are drawn as semicircles between nodes. In this layout, it is important to arrange the nodes to minimize the number of crossings of edge semicircles.

A common type of layout for directed and acyclic graphs is the layered graph, typically layed out from top to bottom or from left to right. The layout begins at the root node or nodes, and increments the layer for each edge between adjacent nodes, as shown in Figure 7.13.

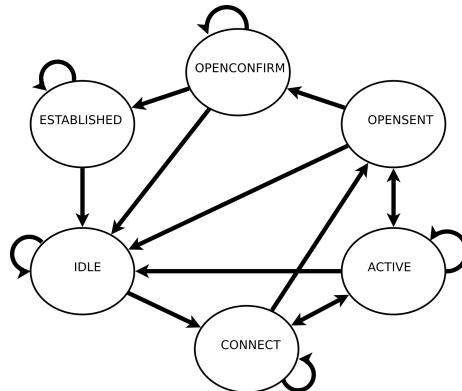
When assessing the quality of the graph layout, a number of considerations are important:

- *Number of crossings of lines or curves.* Such crossings are visually confusing and should be minimized. In fact, this is such an important criterion that graph theory has defined a planar graph as one that can be visualized in two dimensions without any line crossings. Another qual
- *Area of the graph.* Graphs should be drawn in the minimal amount of space while still being easy to read and understand.
- *Symmetries.* Being able to exploit symmetries in the underlying graph data and represent or highlight them in the graphical layout makes the graph visualization easier to understand.



<https://commons.wikimedia.org/wiki/File:SocialNetworkAnalysis.png>

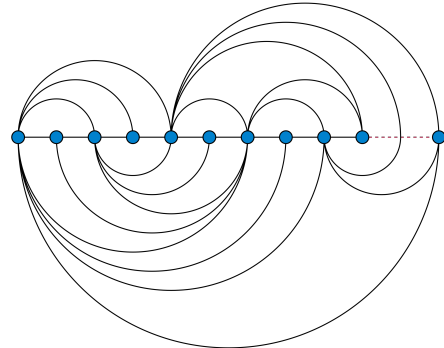
Figure 7.10: Force-directed graph layout example



https://commons.wikimedia.org/wiki/File:BGP_FSM_3.svg

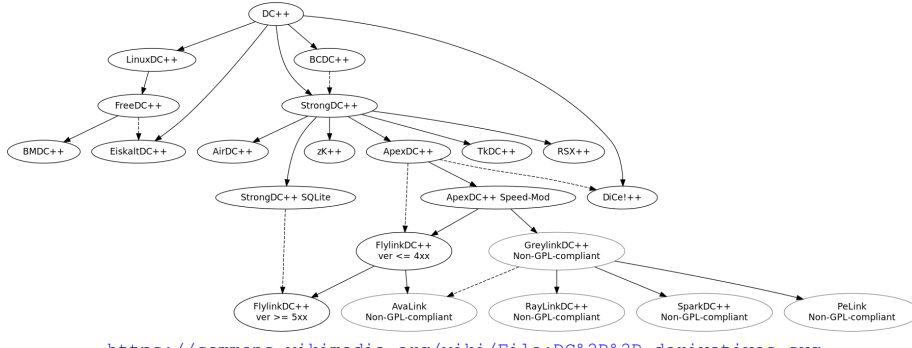
Figure 7.11: Circular graph layout example

- *Shape homogeneity.* A particular problem when using node labels, e.g. for names or node attributes, is the size of the node shape in the graph visualization. It is preferable to maintain equal size, despite differences in label length. Lines should have the lowest or an equal number of bends.
- *Angular resolution.* In graphs with many edges from nodes, it is important to draw lines in such a way that the edges can be clearly differentiated. This is very important in circular layouts, but also plays a large role in other types of graph layouts.



<https://commons.wikimedia.org/wiki/File:Goldner-Harary-linear.svg>

Figure 7.12: Arc graph layout example



https://commons.wikimedia.org/wiki/File:DC%2B%2B_derivatives.svg

Figure 7.13: Layered graph layout example

7.4 Color Palettes

The use of color in data visualization is crucial, serving not only to enhance the visual appeal of a graphic but also to improve its clarity and interpretability. Color choices in data visualizations, determined by the selected color palettes, play a significant role in distinguishing different data points or categories, setting the tone of the presentation (for example, formal versus informal presentations), ensuring accessibility for viewers with color vision deficiencies, and enhancing the overall aesthetic appeal. Desirable characteristics of color palettes are:

- *Range of Values*: Colorful palettes are required when many different values have to be represented and distinguished.
- *Perceptual Uniformity*: The relative perceived differences between colors in the palette should mirror the relative differences in the data values represented by the colors.

- *Robustness to Color Vision Deficiency*: Colour vision deficiency (CFD), colloquially called "color blindness" impacts almost 10% of the population and must therefore be a consideration when choosing color palettes so that the data visualization can be properly perceived and interpreted by everyone.
- *Consistency*: When using multiple plots, their color palette should be the same or at least consistent so as not to cause confusion in interpretation and require less effort for understanding by the reader.
- *Aesthetic Appeal*: Finally, a colour palette should also be "pretty".

Types of Color Palettes

Color palettes can be distinguished by the number of colours they use, and whether the colors span a continuous color space or are a discrete set.

Sequential color palettes

Sequential color palettes, like the one in Figure 7.14a, use a single color and vary the hue or depth of the color. They are best used for data that has an inherent order, as they clearly show progression or gradation. However, they are not suitable for data that lacks a natural ordering. The *monochromatic color palette* is a special case of a sequential palette. This may be suitable when it is likely that the output will be printed on media without the use of color.

Diverging color palettes

Diverging color palettes, like the one in Figure 7.14c, on the other hand, use two colors as anchors and use gradations either through white, as in Figure 7.14c, or through black. They are ideal for emphasizing deviations from a median or mean value, or for highlighting extremes on either side of a critical midpoint. However, these palettes may be misleading if used for data without a meaningful center.

Spectral color palettes

Spectral color palettes, like the one in Figure 7.14d use a variety of different colors without any implicit ordering. They are used to represent discrete categories without inherent ordering, and are useful for differentiating distinct groups of data. The downside is that they can become confusing with too many categories.

Sequential and diverging color palettes may be *discrete*, like the ones shown in Figure 7.14, or *continuous*, while spectral palettes are always discrete.

Color Vision Deficiency

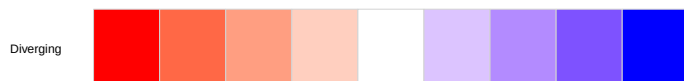
The human eye contains three different types of color receptor cells, called "S-cones" that perceive the color blue, "M-cones" that perceive the color green, and "L-cones" that perceive the color red. Color vision deficiency (CVD) is a biological impairment



(a) Sequential color palette



(b) Monochromatic color palette



(c) Diverging color palette



(d) Spectral color palette

Figure 7.14: Types of Color Palettes

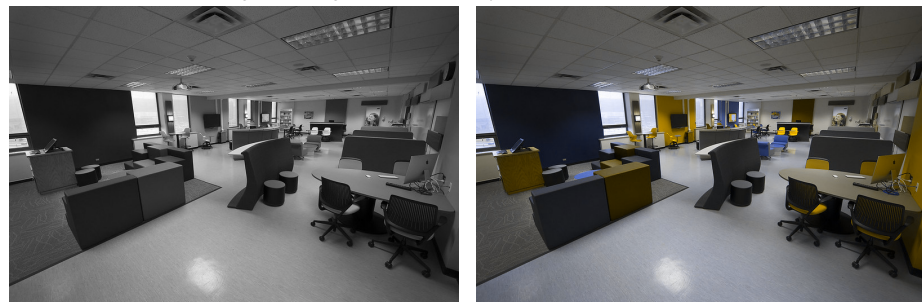
where some color receptor cells in the eye are missing, less frequent, or their function is diminished. In *protanopia*, the S-cones are missing or impaired, in *deutanopia*, the M-cones are missing or impaired, and in *tritanopia*, the L-cones are impaired. When all are missing or non-functional, one speaks of *monochromatism*. CVD is a fairly common disability, afflicting approximately 1 in 12 men and 1 in 200 women, with an overall incidence rate in Canada of more than 5%.

To show the different types of color deficiencies, consider the images in Figure 7.15. Figure 7.15a shows the original image as it is perceived by a person who does not suffer

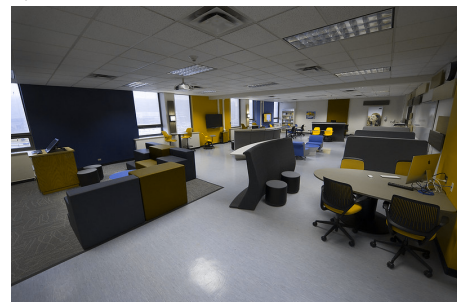


Copyright Memorial University of Newfoundland

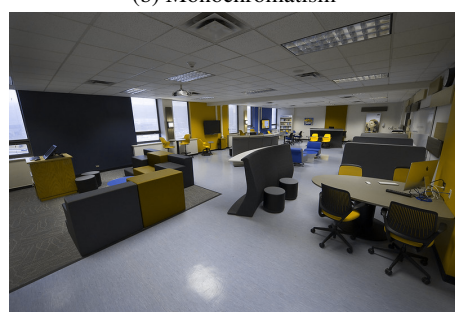
(a) Original Image (MUN Faculty of Education Class Room)



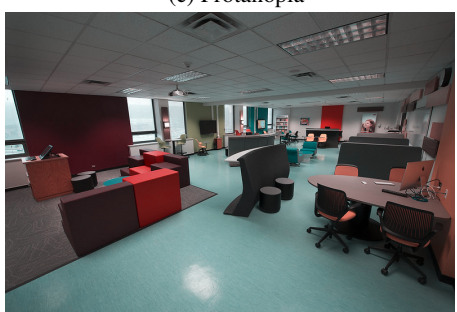
(b) Monochromatism



(c) Protanopia



(d) Deutanopia



(e) Tritanopia

Figure 7.15: Simulated Color Vision Deficiencies

from CVD. The remaining four images show how the photo appears to persons with different types of CVD.

Realizing the prevalence and the effects of CVD means that the color palette that is

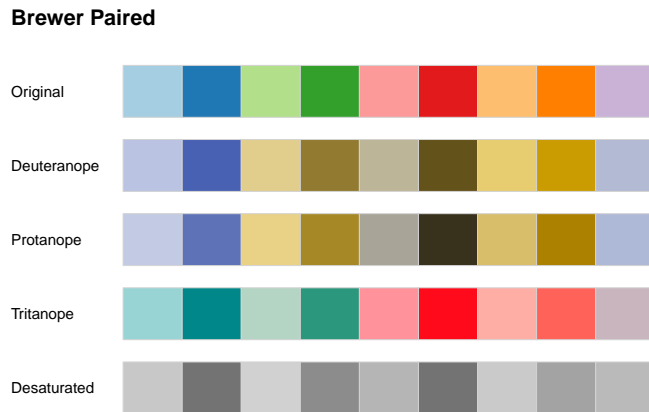


Figure 7.16: Example: Colourbrewer Palette "Paired"

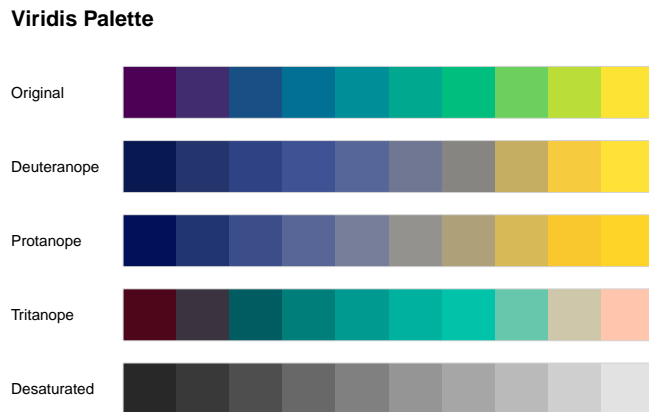


Figure 7.17: Viridis Colour Palette

chosen for data visualization should be interpretable for and lead to the same interpretation even for readers with CVD. For example, the Viridis color palette available in many visualization software packages was designed with CVD readability in mind. Compare the popular "Color Brewer Paired" palette in Figure 7.16 to the Viridis palette in Figure 7.17. The figures show that the Viridis palette is readable and interpretable with any CVD condition, whereas the Paired palette is not because some colors cannot be distinguished under various CVD conditions.

In summary, the thoughtful application of color in data visualization is not merely an artistic decision but a strategic one. It influences how effectively the data is communicated and understood, ensuring that visualizations are not only informative and

accurate, but also inclusive and engaging to a diverse audience.

7.5 Common Types of Plots

Depending on the number of variables to visualize, whether they are discrete or continuous, and the quantitative message to convey, different types of plots may be chosen. While the list of plot types presented here is not comprehensive, and new ways of visualizing data are constantly being invented, these are widely used plot types that are available in most visualization software packages and can be created with little effort.

- Plots for One Variable
 - Continuous
 - * **Area:** Degree of change over time, or relationship of parts to aggregate
 - * **Density, Dot, Frequency, Histogram:** Show frequency distribution of data
 - Discrete
 - * **Bar:** Connections among individual things, compare items of different groups
 - * **Pie:** Relationships of parts to aggregate
- Plots for Two Variables
 - Both Continuous
 - * **Point:** Connections among numeric values, show multiple groups of data
 - * **Lines, Local Regression:** Relationships/correlations among multiple data series or over time
 - * **Text / Label:** Frequency of labels in content/document
 - One Discrete, One Continuous
 - * **Column:** Correlations among things or information changes over time
 - * **Box, Dot, Violin:** Compare distributions between many groups, display spread and skew of data
 - Both Discrete
 - * **Points/Counts:** Magnitude of counts
 - * **Jitter:** Plots of data points
 - Distributions of Two Variables
 - * **Bin2D, Density2D, Hex:** Shows frequency of values over two continuous variables

- Plots for Three Variables
 - Continuous
 - * **Contour, Raster and Tile:** Shows relationships among three data series
- Visualizing Errors and Uncertainty
 - Give a general idea of how precise a value is, or how far a value might be from the true value
 - Typically used to augment a given visualization
 - Common Visualization Styles:
 - * Crossbar
 - * Errorbar
 - * Range (line, point)

7.6 Graphics Libraries and Frameworks

R

The R software system offers several powerful data visualization packages, each with unique features and strengths. Among the most prominent are *ggplot2*, *Plotly* for R, *ggvis*, and *Shiny*, which collectively cater to a wide range of visualization needs.

At the forefront is *ggplot2*, a package based on the Grammar of Graphics, which provides a coherent system for describing and building graphs. Its strength lies in its ability to create complex, multi-layered graphics with a syntax that is both powerful and expressive. *ggplot2*'s approach allows users to build plots layer by layer, making it easier to handle and modify the components of a graphic. Its extensive customization options and the ability to handle a wide variety of graphical forms make it popular for static graphics.

Plotly for R integrates the functionality of the Plotly JavaScript library into R, enabling the creation of interactive, web-based graphs. This package extends the interactive capabilities of R visualizations, allowing users to produce graphics that can be zoomed, panned, and hovered over to reveal additional information. Its integration with R makes it a popular choice for adding an interactive element to data presentations, bridging the gap between static and dynamic visualizations.

ggvis, another package in the R visualization landscape, combines the concepts of *ggplot2* with the interactivity of the web. It is designed to integrate well with R's reactive programming package, *Shiny*, and the *dplyr* package, enabling a smooth workflow for interactive data exploration. *ggvis* focuses on web-based, interactive visualizations, providing a syntax similar to *ggplot2* but with additional capabilities to interactively change the data display and explore data in real-time.

Shiny, distinct from the traditional visualization packages, is an R package for building interactive web applications. It allows users to turn their analyses into interactive web applications without requiring HTML, CSS, or JavaScript knowledge. Shiny applications have the power to not only display complex visualizations but also to interact with the user, making it possible to dynamically change the data, the types of plots, filters, and other aspects of the visualization based on user input. This interactivity makes Shiny particularly useful for creating data dashboards, where users need to explore and interact with data in a flexible manner.

Together, these packages provide R users with a comprehensive toolkit for creating static and interactive visualizations. From detailed and layered static plots with `ggplot2` to dynamic, user-driven applications with Shiny, the R ecosystem enables a vast array of data visualization possibilities, catering to both simple and complex, interactive data exploration and presentation needs.

Python

The Python programming environment also offers a rich landscape of data visualization packages, each tailored to different needs and preferences.

Matplotlib is the foundational library for data visualization in Python, offering a wide array of functionalities to create static, animated, and interactive plots. It is highly customizable and capable of creating virtually any type of chart or graph. The versatility of Matplotlib allows for detailed control over plot elements, but this can also lead to more complex code for intricate visualizations.

Seaborn builds on Matplotlib and simplifies the creation of beautiful, informative statistical graphics. It integrates closely with Pandas, a data manipulation library in Python, and provides a high-level interface for drawing attractive and informative statistical graphics. Seaborn's strength lies in its ability to create complex visualizations like heatmaps, time series, and violin plots with relatively straightforward commands.

Plotnine is inspired by R's `ggplot2` library and brings the Grammar of Graphics to Python. It offers a similar layer-based approach to visualization, making it a familiar choice for users transitioning from R to Python. Plotnine is particularly effective for creating complex, multi-layered graphics with a syntax that emphasizes the declarative nature of the visualization process.

Plotly Express is a high-level interface for the Plotly library, designed to make it easy to create complex, interactive, and beautifully rendered visualizations. It offers a simple syntax for creating a wide variety of chart types and is particularly adept at handling large and complex datasets. Plotly Express's strength lies in its integration with Dash, another Plotly product, for building interactive web applications.

Plotly Graph Objects is the lower-level interface of the Plotly library, providing more granular control over the visualization elements. It's ideal for users who need to create highly customized visualizations or who require fine-tuning beyond what Plotly Express offers.

Plotly Dash is a framework for building interactive web applications with Python (and R and Julia). Dash is unique in its ability to create richly interactive, web-based data visualizations and dashboards without requiring advanced knowledge of web development. It integrates seamlessly with Plotly's suite, allowing for the creation of sophisticated data visualization interfaces.

Bokeh, another prominent Python library, excels in creating interactive and real-time streaming visualizations. It is particularly well-suited for web-based dashboards and applications, offering both simplicity in creating complex interactive plots and the power to handle streaming datasets.

In summary, Python's ecosystem for data visualization is diverse and robust, ranging from Matplotlib's comprehensive capabilities for static plots to the interactive and web-based functionalities of Plotly and Bokeh. Each library offers unique strengths, whether it be in creating complex statistical visualizations, interactive web applications, or real-time data streams, catering to a wide range of data visualization needs and preferences.

JavaScript/Web

JavaScript, being the standard language of web development, boasts several powerful data visualization libraries that are integral for creating interactive and dynamic visualizations on the web. Among these, D3.js, Chart.js, and Google Charts are particularly noteworthy, each with their unique capabilities and strengths.

D3.js stands out as the most sophisticated and flexible JavaScript library for data visualization. Its core strength lies in its ability to bind arbitrary data to a Document Object Model (DOM), and then apply data-driven transformations to the document. D3 allows for extremely detailed and sophisticated visualizations by giving developers direct control over the SVG or HTML output. This level of control enables the creation of complex, interactive, and highly customizable visualizations. However, this power comes with a steep learning curve and can be overkill for simpler visualizations.

Chart.js is a more lightweight and user-friendly alternative, specifically designed for creating simple yet beautiful and interactive charts. It uses HTML5 Canvas for rendering, which makes it efficient in terms of performance. Chart.js supports a variety of chart types, including bar, line, pie, radar, and more, all of which are responsive and mobile-ready by default. Its simplicity and ease of use make it a popular choice for developers who need to implement standard charts quickly and without the complexity of D3.js.

Google Charts provides an even simpler way of incorporating charts into web pages. It offers a wide array of chart types and is particularly known for its integration with other Google services, like Google Spreadsheets. Google Charts is designed to be easy to use, and it handles a lot of the heavy lifting behind the scenes, such as drawing the charts, which makes it an appealing option for users who prefer a more straightforward and less code-intensive approach. The downside is that it offers less customization compared to D3.js and is reliant on external Google services, which might raise privacy concerns or issues with data control.

Each of these libraries serves different needs within the web development and data visualization community. D3.js is ideal for creating complex, interactive visualizations where control and customization are paramount. Chart.js offers a balance between simplicity and functionality, suitable for standard web-based charts. Google Charts, with its ease of use and integration with Google products, is excellent for straightforward visualizations where ease of implementation is a priority. The choice among these libraries largely depends on the specific requirements of the project, the complexity of the visualizations needed, and the developer's proficiency with JavaScript and web technologies.

7.7 Mapping Data to Plot Elements

Creating a basic visualization in two dimensions, such as bar chart, a line chart, or a bubble chart, means that data elements or data series must be mapped to visualization elements. This is the core of the visualization task, and the most fundamental choice the data analyst has to make. Table 7.1 shows plot elements that data variables can be mapped to. In principle, a different data variable can be mapped to each of these, resulting in potentially being able to represent more than a dozen variables in one diagram. However, in practice, the number of concurrent variables to represent should be limited to no more than 3, in order for the visualization to remain interpretable and not to require too much cognitive effort on the part of the reader.

X, Y, Z axes
Colour (of points, lines, areas, shapes)
Transparency ("alpha")
Patterns (within areas, shapes)
Size, Weight/Width (of points and lines)
Shape, Style (of points and lines)

Table 7.1: Plot elements that can be mapped to data variables

7.8 Visualization in R using ggplot2

This section provides an introduction to data visualization using the ggplot library in R. The example dataset for this section is the Fuel Consumption Ratings for battery electric vehicles, provided the Government of Canada through its Open Government Portal³. At the time of writing, the dataset was last updated on October 10, 2023. The dataset contains the variables shown in Table 7.2.

Reading and preprocessing the data is straightforward in R, shown in the following code block:

³<https://open.canada.ca/data/en/dataset/98f1a129-f628-4ce4-b24d-6f16bf24dd64>

Column	Data Type	Definition
Make	Discrete	Manufacturer
Model	Discrete	Model name
Year	Numeric	Model year
Category	Discrete	Small, Midsize, Large, Pickup, SUV, Station Wagon, etc.
City	Numeric	Consumption in l/100km equiv.
Hwy	Numeric	Consumption in l/100km equiv.
Comb	Numeric	Consumption in l/100km equiv.
Range	Numeric	Driving range in km

Table 7.2: Fuel efficiency data set variables

```
# Import libraries
library(tidyverse)
# Read CSV
e <- read.csv('https://evermann.ca/busi4720/fuel.csv')
# Pre-process for data types
e$Year <- as.numeric(e$Year)
e$Category <- as.factor(e$Category)
e$Fuel <- as.factor(e$Fuel)
e$City <- as.numeric(e$City)
e$Hwy <- as.numeric(e$Hwy)
e$Comb <- as.numeric(e$Comb)
e$Range <- as.numeric(e$Range)
e$Annual <- as.numeric(e$Annual)
e.clean <- e
```

Next, load the required graphics libraries. A number of extensions to the core ggplot2 library have been developed to provide additional capabilities, such as radar plots, pattern fills, providing more control over scales and axes, etc.

```
library(ggplot2)
library(ggpattern)
library(ggstream)
library(ggsci)
library(scales)
library(ggrepel)
library(ggradar)
```

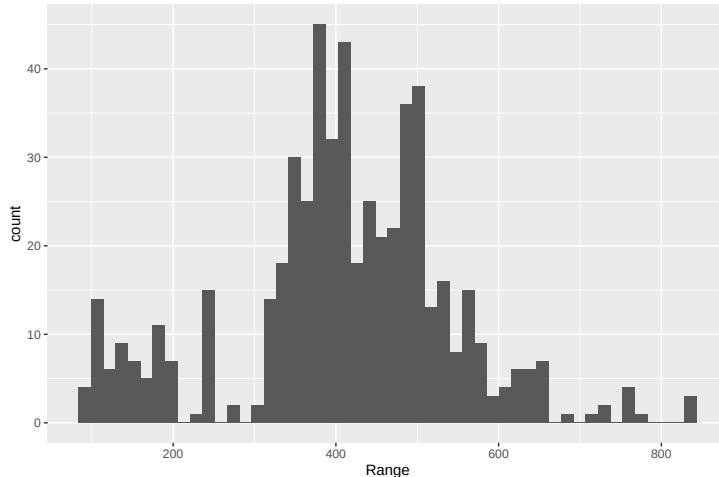
The core `ggplot()` function can be used in a dplyr pipeline and accepts the processed data tibble. The core argument to `ggplot()` is the "aesthetic" that maps plot elements to data variables. The actual plots themselves are then added through the use of various "geoms". Such geoms represent commonly used plot types. The geoms "inherit" the aesthetic specified in `ggplot()` and can add to it by including more variables mapped

to different plot elements. More than one geom can be added to a plot, allowing the analyst to overlay plot types or combine plots for multiple data series or data sets. The final graph can be saved in a variety of different image formats.

The first example below introduces the histogram geom. Histograms show the count of values in a certain range. The `ggplot()` function's aesthetic maps the "Range" variable of the tibble to the x axis of the plot. The argument to the `geom_histogram()` function indicates that 50 bins should be formed, i.e. the data is divided in 50 separate regions for counting and plotting. The `ggsave()` function saves the last plot in a file with the specified height and width.

```
e.clean |>
  ggplot(aes(x=Range)) +
  geom_histogram(bins=50)

ggsave("histogram.pdf",
       height=5, width=7.5, units='in')
```



A density plot using the `geom_density()` function, is similar to a histogram in that it indicates the frequency of values. However, a density plot shows a continuous probability distribution of the data values, and as such is limited in range between 0 and 1.

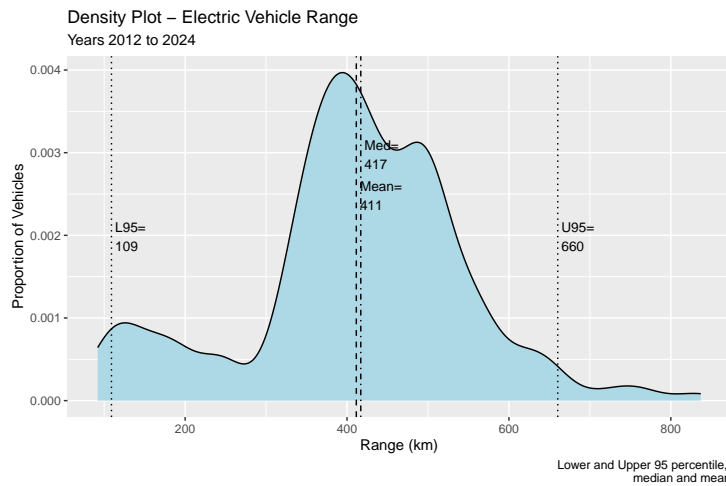
The example below adds a number of elements to the basic density plot. The function `labs()` allows specification of labels for all plot elements. The `geom_vline()` geoms add vertical lines. Note that these geoms do not receive the data from the pipe, but the data is specified using the `data` argument. The aesthetics of a vertical line map the x axis intercept to a data variable. Different line types are used for the different lines. The `annotate()` function adds text annotations to the plot. Each annotation prints a label at a set of x and y coordinates in the plot, with a specific size and horizontal justification. For example, `hjust=0` means the text is left-justified. Consult

the documentation for further details.

```
# Prepare summary statistics

mean_v <- e.clean |>
  summarize(mean_v = mean(Range),
            median_v = median(Range),
            lower95=quantile(Range, .025),
            upper95=quantile(Range, .975),
            maxdensity = max(density(Range)$y))

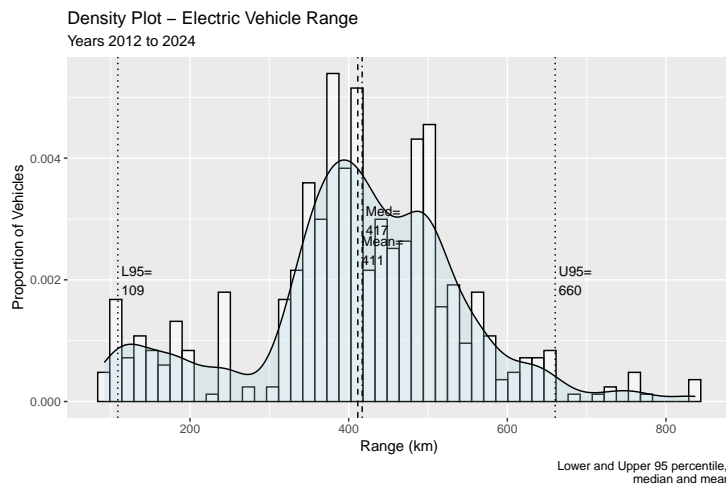
e.clean |>
  ggplot(aes(Range)) +
  geom_density(kernel='gaussian',
               fill='lightblue') +
  labs(x = 'Range (km)',
       y = 'Proportion of Vehicles',
       title='Density Plot - Electric Vehicle Range',
       subtitle='Years 2012 to 2024',
       caption='Lower and Upper 95 percentile,
                 median and mean') +
  geom_vline(data=mean_v,
             aes(xintercept=mean_v),
             linetype='dashed') +
  geom_vline(data=mean_v,
             aes(xintercept=median_v),
             linetype='dotdash') +
  geom_vline(data=mean_v,
             aes(xintercept=lower95),
             linetype='dotted') +
  geom_vline(data=mean_v,
             aes(xintercept=upper95),
             linetype='dotted') +
  annotate('text',
           label=paste(' L95=\n ', round(mean_v$lower95), sep=' '),
           x = mean_v$lower95, y = mean_v$maxdensity/2,
           size=3.5, hjust=0) +
  annotate('text',
           label=paste(' Med=\n ', round(mean_v$median_v), sep=' '),
           x = mean_v$median_v, y = mean_v$maxdensity*3/4,
           size=3.5, hjust=0) +
  annotate('text',
           label=paste(' Mean=\n ', round(mean_v$mean_v), sep=' '),
           x = mean_v$mean_v, y = mean_v$maxdensity*5/8,
           size=3.5, hjust=0) +
  annotate('text',
           label=paste(' U95=\n ', round(mean_v$upper95), sep=' '),
           x = mean_v$upper95, y = mean_v$maxdensity/2,
           size=3.5, hjust=0)
```



The next example shows how histograms and density plots can be combined. The R code fragment below indicates only the relevant changes to the previous example.

```
...
  geom_histogram(aes(y=..density..), bins=50,
                 alpha=0.5, fill='white', color='black', ) +
  geom_density(kernel='gaussian',
               alpha=0.25, fill='lightblue') +
...

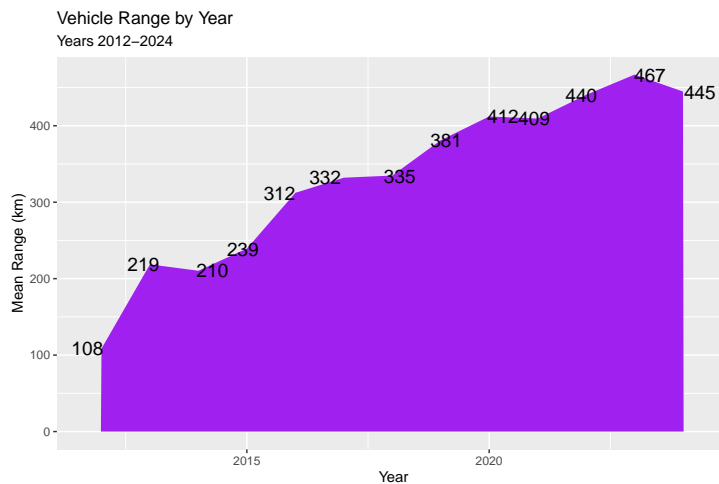
```



An area plot is essentially a line plot where the area under the line filled. However, in contrast to a line plot, when plotting multiple data variables in an area plot, the area plot is cumulative, that is, data are stacked on top of each other. The following example first uses dplyr functions to compute a summary statistic, and then pipes the

result into the `ggplot()` function. `geom_text()` is another way to add annotations to the data. This geom uses its own aesthetic in the example below. Note that `position='jitter'` indicates that overlapping points (i.e. points with the same data values) should be randomly moved a little bit to show them separately.

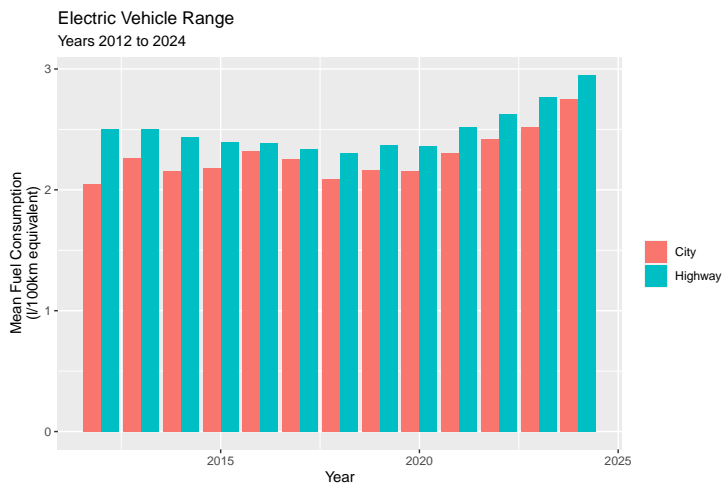
```
e.clean %>%
  group_by(Year) %>%
  summarize(meanRange = mean(Range)) %>%
  ungroup() %>%
  ggplot(aes(Year, meanRange)) +
  geom_area(fill='purple') +
  geom_text(aes(label=round(meanRange)),
            size=5, position='jitter') +
  labs(x='Year', y='Mean Range (km)',
       title='Vehicle Range by Year',
       subtitle='Years 2012-2024')
```



The next example shows a column chart. Again, dplyr functions are used to create suitable summary statistics to plot, and the summarized data is then piped to `ggplot()`. The variable "metric" is mapped to the "fill" element of the plot, that is the color with which columns are filled. The `position='dodge'` argument to the `geom_col()` function indicates that columns are located next to each other, instead of being stacked on top of each other.

This example also shows customization of the scales. Here, the fill scale (that is, the colour) is customized, first by specifying a colour palette, and then by providing labels for the different categories. The legend is automatically added to the right of the column plot.

```
e.clean %>%
  group_by(Year) %>%
  summarize(meanCity = mean(City), meanHwy = mean(Hwy)) %>%
  ungroup() %>%
  pivot_longer(cols=c('meanCity', 'meanHwy'),
               names_to='metric',
               values_to='consumption') %>%
  ggplot(aes(Year, consumption, fill=metric)) +
  geom_col(position='dodge') +
  scale_fill_brewer(palette="Paired") +
  scale_fill_discrete(labels=c("City", "Highway")) +
  labs(x = 'Year',
       y='Mean Fuel Consumption\n(l/100km equivalent)',
       fill='',
       title='Electric Vehicle Range',
       subtitle='Years 2012 to 2024')
```



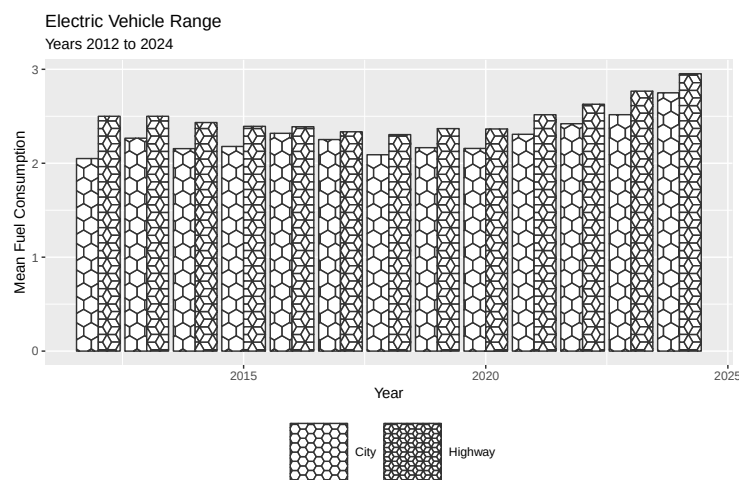
When it is clear that a plot is likely to be printed in black and white, it may be useful to omit the use of colours and instead use different fill patterns. They are provided by the `ggpattern` package that provides the `geom_col_pattern` geom. As shown in the code below, the aesthetics for this geom can map data values to different aspects of a fill pattern, such as the pattern type and the pattern angle.

This example also customizes the scale for the "pattern" geom to provide values for the different pattern types and labels for the two data series. Note that more than the required two values for patterns are listed in the example to showcase the different options provided by the `ggpattern` package. Additionally, the `guides()` function is used to omit the legend for the pattern angle (because the same data is mapped to pattern type and pattern angle) and the `theme()` function customizes the layout of the legend.

```

e.clean %>%
  group_by(Year) %>%
  summarize(meanCity = mean(City),
            meanHwy = mean(Hwy)) %>%
  ungroup() %>%
  pivot_longer(
    cols=c('meanCity', 'meanHwy'),
    names_to='metric',
    values_to='consumption') %>%
  ggplot(aes(Year, consumption)) +
  geom_col_pattern(
    aes(pattern_type=metric, pattern_angle=metric),
    pattern='polygon_tiling',
    pattern_fill='white',
    pattern_scale=0.5,
    position='dodge',
    pattern_key_scale_factor=0.4) +
  scale_pattern_type_manual(
    values = c('hexagonal', 'rhombille', 'pythagorean',
              'truncated_square', 'rhombitrihexagonal',
              'truncated_trihexagonal'),
    labels=c("City", "Highway")) +
  labs(x = 'Year', y='Mean Fuel Consumption',
       pattern_type='',
       title='Electric Vehicle Range',
       subtitle='Years 2012 to 2024') +
  guides(pattern_angle=FALSE,
         pattern_type=guide_legend(nrow=1)) +
  theme(legend.key.size=unit(1.5, 'cm'),
        legend.position='bottom')

```



A box plot, also known as a box-and-whisker plot, is a way of displaying the distribution of data based on 5 summary statistics: the minimum, first quartile (Q1), median, third quartile (Q3), and the maximum. A box plot provides a visual summary of the

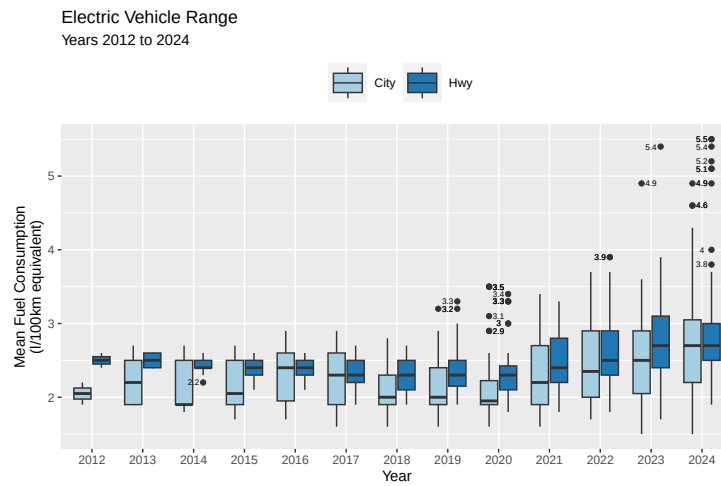
spread, central tendency, and symmetry of the data. Boxplots contain the following elements:

- *The Box:* The bottom and top edges of the box represent the first quartile (Q1, the 25th percentile) and the third quartile (Q3, the 75th percentile), respectively. The box therefore describes the interquartile range (IQR), i.e. the distance between the first and third quartiles.
- *The Median:* Inside the box, there is usually a line that denotes the median (the 50th percentile) of the dataset. By comparing the placement of the median line to the extent of the first and third quartiles, one can judge whether the data is skewed.
- *Whiskers:* Extending from the box are lines called whiskers. One common method is to extend the whiskers to the furthest data point within 1.5 times the IQR from the quartiles. This means that the lower whisker extends to the smallest data point greater than $Q1 - 1.5 * \text{IQR}$ and the upper whisker extends to the largest data point less than $Q3 + 1.5 * \text{IQR}$.
- *Outliers:* Data points that fall outside of the whiskers are often considered outliers and may be plotted as individual points.

Box plots are particularly useful for displaying the distribution of data, comparing multiple distributions, and identifying outliers.

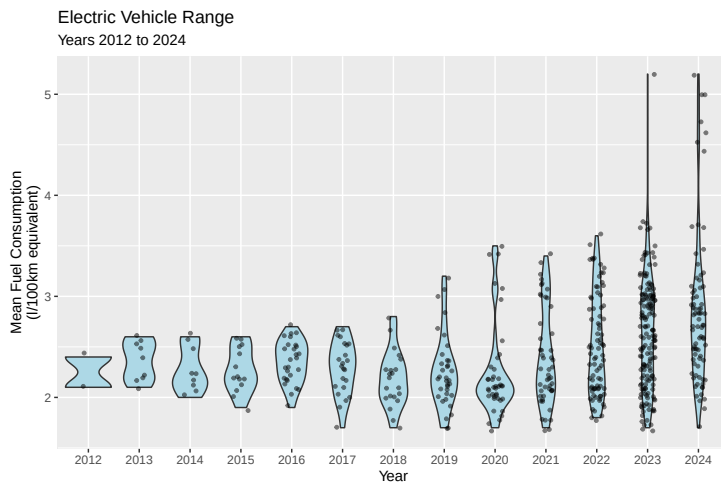
The following example introduces the `stat_summary()` function to add a summary statistic in the form of a text label to the plot. The label is computed by the `stat(y)` function, which in turn calls `fun.y` specified within the `stat_summary()` function itself. The function as used in this example determines how to label the outlier points in the box plot, using the `text` geom and its label.

```
e.clean %>%
  pivot_longer(cols=c('City', 'Hwy'),
               names_to='metric',
               values_to='consumption') %>%
  ggplot(aes(x=as.factor(Year), y=consumption, fill=metric)) +
  geom_boxplot() +
  stat_summary(
    aes(label = round(stat(y), 1)),
    geom = "text",
    size=2,
    fun.y = function(y) {
      o<-boxplot.stats(y)$out;
      if(length(o)==0) NA else o}) +
  scale_fill_brewer(palette="Paired") +
  labs(x = 'Year',
       y='Mean Fuel Consumption\n(l/100km equivalent)',
       fill='',
       title='Electric Vehicle Range',
       subtitle='Years 2012 to 2024') +
  theme(legend.key.size=unit(1, 'cm'),
        legend.position='top')
```



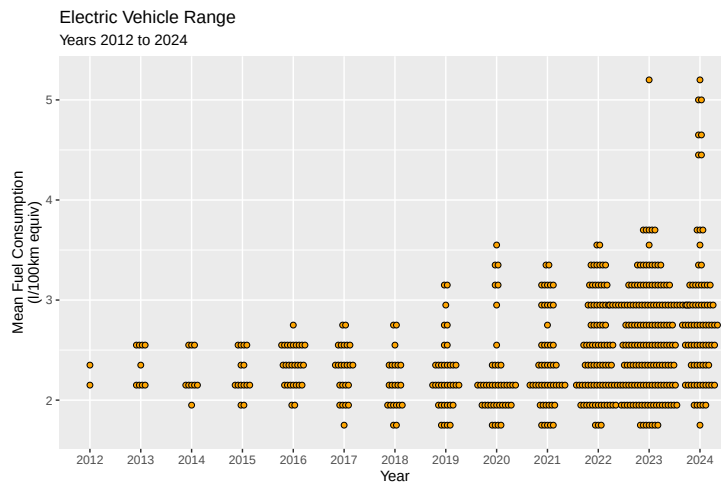
Violin plots are another way to visualize the spread and distribution of the data. Their width is determined by the frequency/distribution of data points. The following example introduces the `geom_violin` geom and the `geom_jitter` geom. As the name suggests, the jitter geom plots and "jitters" the data points, by moving them slightly to avoid overlap. The arguments provided to `geom_jitter()` set the width, the color and size of the points, the fill color and the transparency level ("alpha"). The plot indicates the distribution of data, which is reinforced by the visual "density" of the individual data points in the plot.

```
e.clean %>%
  ggplot(aes(x=as.factor(Year), y=Comb)) +
  geom_violin(fill='lightblue') +
  geom_jitter(width=0.15, color='black',
              size=1, fill=NA, alpha=0.5) +
  scale_fill_brewer(palette="Paired") +
  labs(x = 'Year',
       y='Mean Fuel Consumption\n(l/100km equivalent)',
       fill='',
       title='Electric Vehicle Range',
       subtitle='Years 2012 to 2024')
```



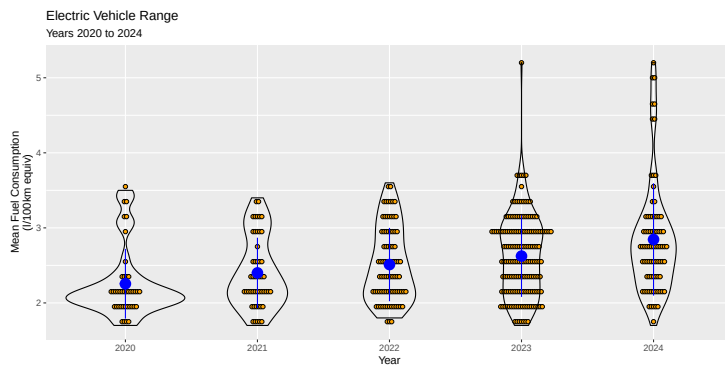
A dot plot is useful for showing individual data points. In this example, the data are binned along the y axis (that is, by combined fuel economy). The stack ratio determines the horizontal separation of points.

```
e.clean %>%
  ggplot(aes(x=as.factor(Year), y=Comb)) +
  geom_dotplot(binaxis='y',
               stackdir='center',
               stackratio=0.5,
               binpositions='all',
               dotsize=0.5,
               color='black',
               fill='orange') +
  scale_fill_brewer(palette="Paired") +
  labs(x = 'Year',
       y='Mean Fuel Consumption\n(l/100km equiv)',
       fill='',
       title='Electric Vehicle Range',
       subtitle='Years 2012 to 2024')
```



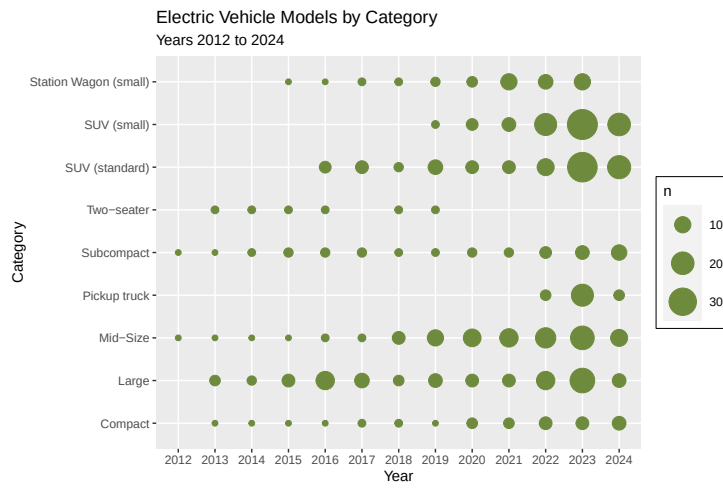
Instead of using a jitter plot with a violin plot, it is sometimes better to combine a violin plot with a dot plot, as in the following example.

```
e.clean %>%
  filter(Year > 2019) %>%
  ggplot(aes(x=as.factor(Year), y=Comb)) +
  geom_dotplot(binaxis='y',
                stackdir='center', stackratio=0.5,
                binpositions='all', dotsize=0.5,
                color='black', fill='orange') +
  geom_violin(color='black', fill=NA) +
  stat_summary(fun.data=mean_sdl,
               fun.args=list(mult=1),
               size=1, color='blue',
               geom="pointrange") +
  scale_fill_brewer(palette="Paired") +
  labs(x = 'Year',
       y='Mean Fuel Consumption\n(l/100km equiv)',
       fill='',
       title='Electric Vehicle Range',
       subtitle='Years 2020 to 2024') +
  theme(legend.position='none')
```



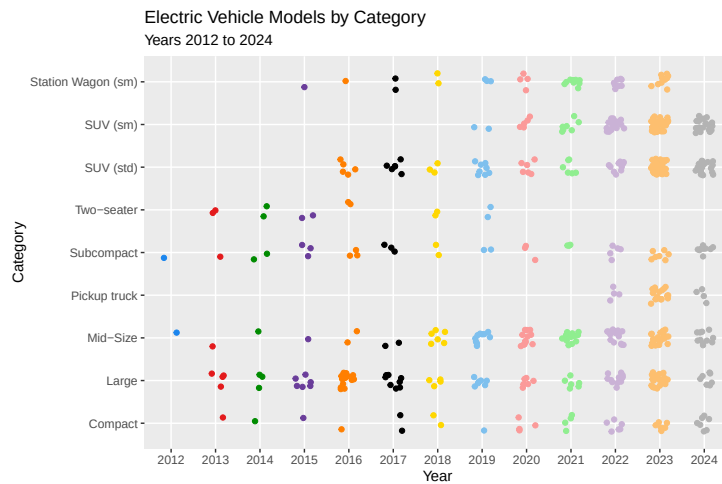
A count plot is useful to show the count of data values as the size of a point. In the following example, the point size is determined by the count of values in each combination of "Year" and "Category". All points have the same color, and the area is scaled to a maximum size of 10 using 6 different sizes. Additionally, this examples shows further customization of the plot legend using the `theme()` function, by surrounding the legend with a black rectangle without fill (transparent). The `guides()` function omits a legend for the color.

```
e.clean %>%
  ggplot(aes(as.factor(Year), as.factor(Category))) +
  geom_count(color='darkolivegreen4') +
  scale_size_area(max_size=10, n.breaks=6) +
  scale_color_brewer(palette="Paired") +
  scale_y_discrete(
    labels=c('Compact', 'Large', 'Mid-Size', 'Pickup truck',
           'Subcompact', 'Two-seater', 'SUV (standard)',
           'SUV (small)', 'Station Wagon (small)')) +
  guides(color=FALSE) +
  labs(x = 'Year',
       y='Category',
       fill='',
       title='Electric Vehicle Models by Category',
       subtitle='Years 2012 to 2024') +
  theme(legend.background=element_blank(),
        legend.box.background=element_rect(color='black', fill=NA),
        legend.key.size=unit(1, 'cm'))
```



A similar effect can be achieved with jitter plot, where the size of the points "cloud" indicates is used analogous to the size of the point. Visually, the following plot achieves a similar goal as the previous dot plot. Here, the same variable is mapped to both the x axis as well as the color element, but the `guides` function omits a legend for the colour element.

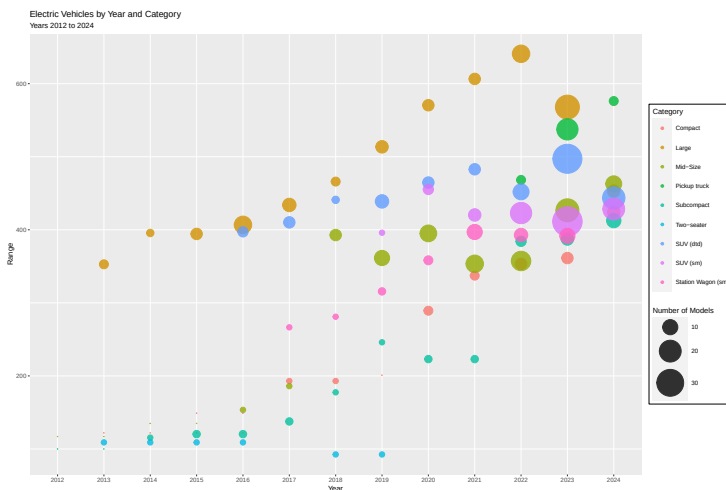
```
e.clean %>%
  ggplot(aes(x=as.factor(Year),
             y=as.factor(Category),
             color=as.factor(Year))) +
  geom_jitter(width=0.2, height=0.2) +
  scale_color_manual(values=c25) +
  scale_y_discrete(
    labels=c('Compact', 'Large', 'Mid-Size',
            'Pickup truck', 'Subcompact',
            'Two-seater', 'SUV (std)',
            'SUV (sm)', 'Station Wagon (sm)')) +
  guides(color=FALSE) +
  labs(x = 'Year',
       y='Category',
       fill='Make',
       title='Electric Vehicle Models by Category',
       subtitle='Years 2012 to 2024')
```



A points plot, sometimes called a bubble chart, generalizes the count plot. Whereas the count plot uses the number of data values to determine the size of the point, the points plot allows one to provide an explicit mapping for the point size. However, the following example also maps the size of the point to the count of values by "Year" and "Category", while the colour is mapped to the "Category" variable.

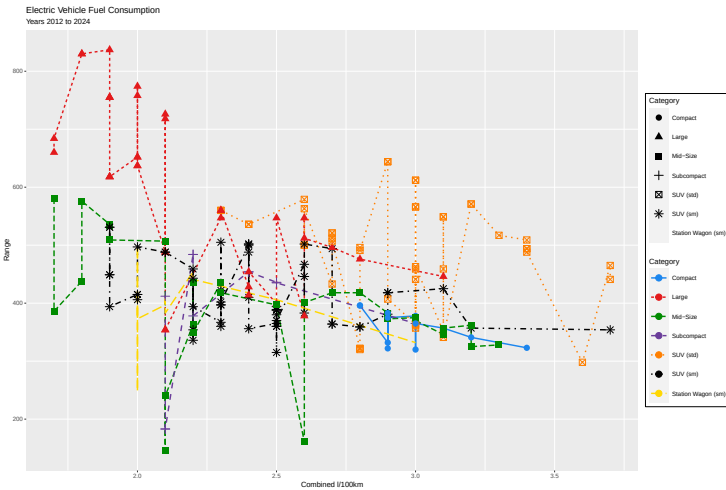
The point size scale is continuous in the range from 0 to 20, the color scale is set to the "tron" colour palette, while the y axis is continuous. The colous are mapped to specific labels for displaying in the legend. Note that the legend contains information both for the size as well as the colour of the points.

```
e.clean %>%
  group_by(Year, Category) %>%
  summarize(totalcount=n(), meanRange=mean(Range)) %>%
  ungroup() %>%
  ggplot(aes(x=as.factor(Year), y=meanRange,
             size=totalcount, color=Category)) +
  geom_point(alpha=0.8) +
  scale_size_continuous(range=c(0, 20)) +
  scale_color_tron() +
  scale_y_continuous(labels=scales::comma) +
  scale_color_discrete(
    labels=c('Compact', 'Large', 'Mid-Size', 'Pickup truck',
           'Subcompact', 'Two-seater', 'SUV (std)',
           'SUV (sm)', 'Station Wagon (sm)')) +
  labs(x = 'Year', y='Range',
       fill='Make', size='Number of Models',
       title='Electric Vehicles by Year and Category',
       subtitle='Years 2012 to 2024', ) +
  guides(color=guide_legend(position='right'),
         size=guide_legend(position='right')) +
  theme(legend.background=element_rect(color='black', fill=NA),
        legend.box.background=element_rect(color='black', fill=NA),
        legend.key.size=unit(1, 'cm'))
```



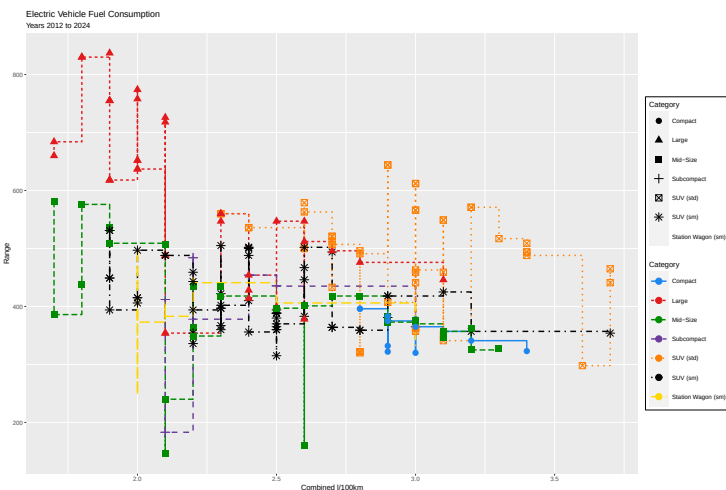
The next example uses two geoms, `geom_line()` to show a line plot and `geom_point()` to also include the data points themselves. While visually not very informative in this case, the example illustrates an aesthetic that maps variables to five different plot elements. However, the same variable "Category" is here mapped to three different plot elements, the colour (of both points and lines), the shape of a point, and the style or type of the line. The code R fragment below omits specification of labels and theme information for the legend, which may be assumed similar to the above example.

```
e.clean %>%
  filter(Year >= 2022 & Year <= 2023) %>%
  filter(Comb <= 4) %>%
  filter(Category != 'PL') %>%
  filter(Category != 'T') %>%
  ggplot(aes(Comb, Range,
             color=Category,
             shape=Category,
             linetype=Category)) +
  geom_line(size=1) +
  geom_point(size=4) +
  scale_color_manual(values=c25,
                     labels=c('Compact', 'Large', 'Mid-Size',
                             'Subcompact', 'SUV (std)',
                             'SUV (sm)', 'Station Wagon (sm)')) +
  scale_linetype(
    labels=c('Compact', 'Large', 'Mid-Size',
            'Subcompact', 'SUV (std)',
            'SUV (sm)', 'Station Wagon (sm)')) +
  scale_shape(
    labels=c('Compact', 'Large', 'Mid-Size',
            'Subcompact', 'SUV (std)',
            'SUV (small)', 'Station Wagon (sm)')) +
  ...
```



To add "steps" to the line, one can use the `geom_step()` instead of `geom_line()`, as in the following example.

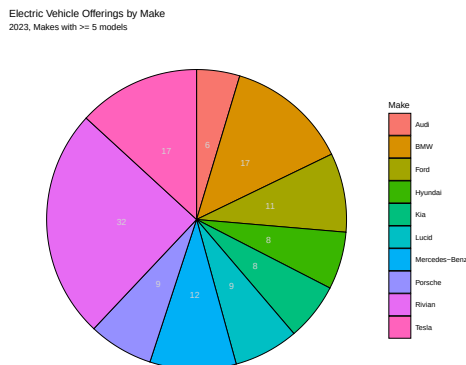
```
...  
  geom_step(size=1) +  
...
```



A pie chart is produced in ggplot2 by taking a stacked bar chart, and "bending" it by plotting on a polar coordinate system. The following example uses the `coord_polar()` function to specify a coordinate system where the "y" axis is mapped to the angle of rotation, `direction=-1` indicates clock-wise rotation and `start=0` indicates to begin the chart at the top of the "pie". The `geom_text()` geom is used to specify labels and compute their position in the pie chart. It provides its own aesthetic

for the label's color and position. Note that it assumes a stacked bar chart so that `position_stack(vjust=0.5)` positions the label vertically in the center of the area. When plotted in the polar coordinate system, this translates to the label centered in the pie slice.

```
e.clean %>%
  filter(Year==2023) %>%
  group_by(Make) %>% summarize(totalcount = n()) %>%
  filter(totalcount >= 5) %>%
  ungroup() %>%
  ggplot(aes(x='', y=totalcount, fill=Make)) +
  geom_bar(stat='identity',
            color='black', size=0.25, width=1) +
  coord_polar('y', direction=-1, start=0) +
  geom_text(aes(
    label = ifelse(totalcount >= 5, totalcount, '')),
            color='lightgrey',
            position = position_stack(vjust=0.5)) +
  scale_y_continuous(labels=NULL) +
  scale_color_brewer(palette="Paired") +
  labs(x = '', y = '', fill='Make',
       title='Electric Vehicle Offerings by Make',
       subtitle='2023, Makes with >= 5 models') +
  theme_void() +
  theme(legend.key.size=unit(1, 'cm'))
```

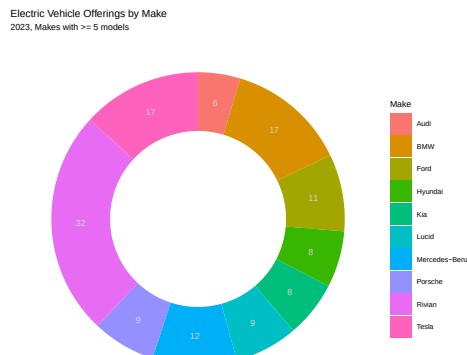


A donut chart is simply a pie chart with a hole in the center. As the pie chart in ggplot2 is a "bent" bar chart, the hole is achieved by adding "whitespace" to the right of the stacked bars (that is, by moving the x axis limits), which will end up getting "bent" into the hole in the center (recall that the `coord_polar()` function bends clock-wise).

```

holesize <- 2
.....
ggplot(aes(x=holesize, y=totalcount, fill=Make)) +
  geom_col() +
  xlim(c(0.2, holesize+0.5)) +
  .....

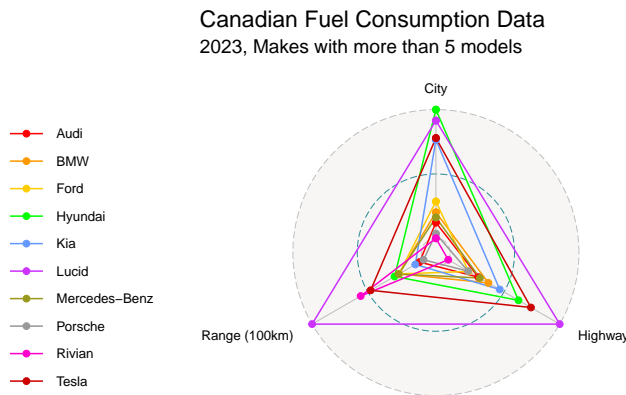
```



A radar plot, sometimes called a spiderweb plot, is useful to show a comparison of different objects on a range of variables. In R, the radar plot is produced by its own library, `ggradar`. In contrast to `ggplot`, `ggradar` requires its data in "wide" format, that is, rather than a single column that provides values for different categories, the values for each category must be provided in their own column.

The following example computes some summary statistics, and scales the resulting variables to a range between 0 and 1, i.e. it standardizes them using the `mutate_at()` function. The radar plot does not require an aesthetic specification, as it is based on the number of columns in the data frame or tibble provided from the pipe.

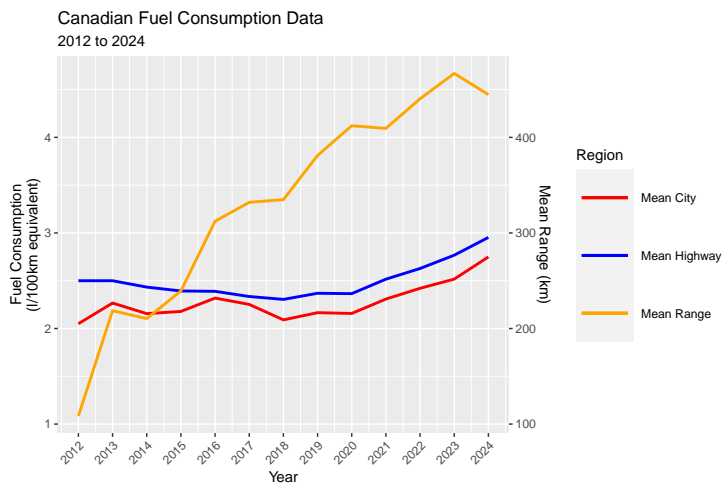
```
e.clean %>%
  filter(Year == 2023) %>% group_by(Make) %>%
  summarize(meanCity = 1/mean(City),
            meanHwy = 1/mean(Hwy),
            meanRange = mean(Range)/100,
            nModels = n()) %>%
  filter(nModels >= 5) %>% ungroup() %>%
  select(-nModels) %>%
  mutate_at(vars(-Make), rescale) %>%
  ggRADAR(axis.labels =
              c('City', 'Highway', 'Range (100km)'),
              values.radar='',
              group.line.width=0.75,
              group.point.size=3) +
  scale_color_UCSCGB() +
  labs(x = '', y = '',
       fill='Make',
       title='Canadian Fuel Consumption Data',
       subtitle='2023, Makes with more than 5 models')
```



Sometimes, it is useful to compare trends of variables that use different scales. However, beware of the potential for misuse; that is, it is easy to visually suggest correlations where none exist. The following example includes three line plots (`geom_line()`) and specifies a secondary y axis using `sec.axis=sec_axis(...)`. In ggplot2, the secondary axis cannot be arbitrary but must be a scaled version of the primary axis. In this example, it is scaled by one hundred using the formula $y_2 = y_1 / 100$.

. *100. Accordingly the data is provided by dividing by a hundred using `mutate()`.

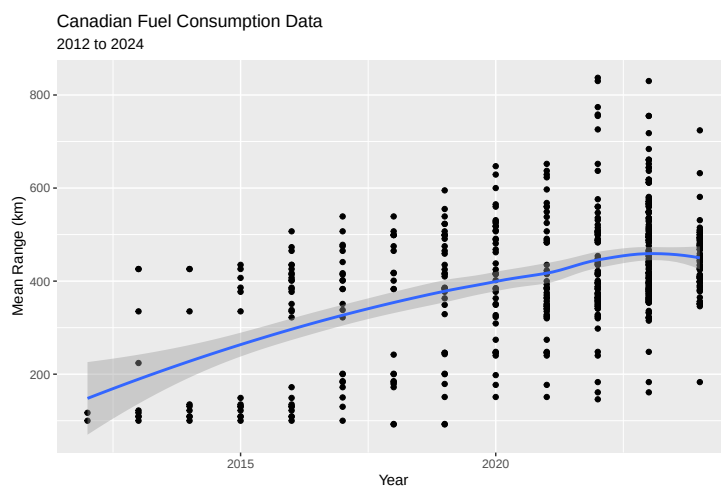
```
e.clean %>%
  group_by(Year) %>%
  summarize(meanCity = mean(City),
            meanHwy = mean(Hwy),
            meanRange = mean(Range)) %>%
  ungroup() %>%
  mutate(meanRange2 = meanRange/100) %>%
  ggplot(aes(x=Year)) +
  scale_color_manual(name='Region',
                     values=c('Mean City' = 'red',
                             'Mean Highway' = 'blue',
                             'Mean Range' = 'orange')) +
  geom_line(aes(y=meanCity, color='Mean City')) +
  geom_line(aes(y=meanHwy, color='Mean Highway')) +
  geom_line(aes(y=meanRange2, color='Mean Range')) +
  scale_y_continuous(labels=scales::comma,
                      name="Fuel Consumption\n(l/100km equiv)",
                      sec.axis=sec_axis(~ .*100,
                                       labels=scales::comma,
                                       name="Mean Range (km)") +
  scale_x_continuous(breaks=seq(from=2012,to=2024,by=1)) +
  labs(x = 'Year', color='',
       title='Canadian Fuel Consumption Data',
       subtitle='2012 to 2024') +
  theme(legend.key.size=unit(1.5, 'cm'),
        axis.text.x = element_text(angle=45, hjust=1))
```



So called "trendlines" can be added to plots easily with the `geom_smooth` geom. Different options to compute the trendlines exist, but the most frequently used one is the local polynomial regression, where the slope of the line is determined by a regression that uses data points in the vicinity of the line, weighted by their proximity. As with any regression, there is uncertainty around the estimated slope parameter (standard

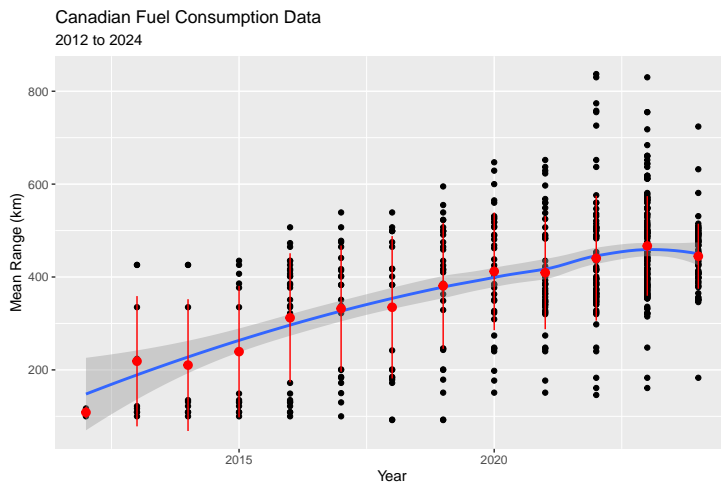
deviation or variance) and this uncertainty can be visualized as well, as shown in the plot below by the gray area around the trendline. Note that the uncertainty is greater in areas where there are fewer data points, as would be expected.

```
e.clean %>%
  ggplot(aes(Year, Range)) +
  geom_point() +
  geom_smooth() +
  scale_y_continuous(labels=scales::comma) +
  labs(x = 'Year', color='', y = 'Mean Range (km)',
       title='Canadian Fuel Consumption Data',
       subtitle='2012 to 2024')
```

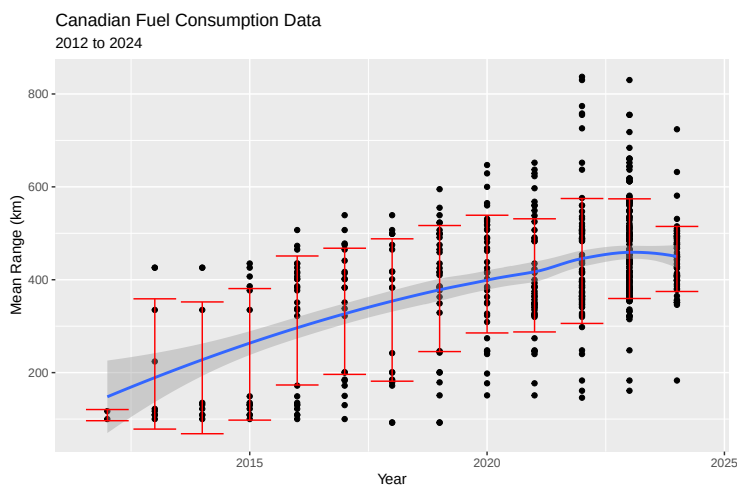


The next three examples augment the previous plot with indicators for the variability or spread of the data. First, a red point range indication is overlayed (`geom="pointrange"`) using the `stat_summary()` function. This shows mean and standard deviation. Another way to visualize variability is with the error bars or cross bars. All show the same information, but in different ways:

```
e.clean %>%
  ggplot(aes(Year, Range)) +
  geom_point() +
  geom_smooth() +
  stat_summary(
    fun.data=mean_sdl,
    fun.args=list(mult=1),
    color='red',
    geom="pointrange") +
  ...
```

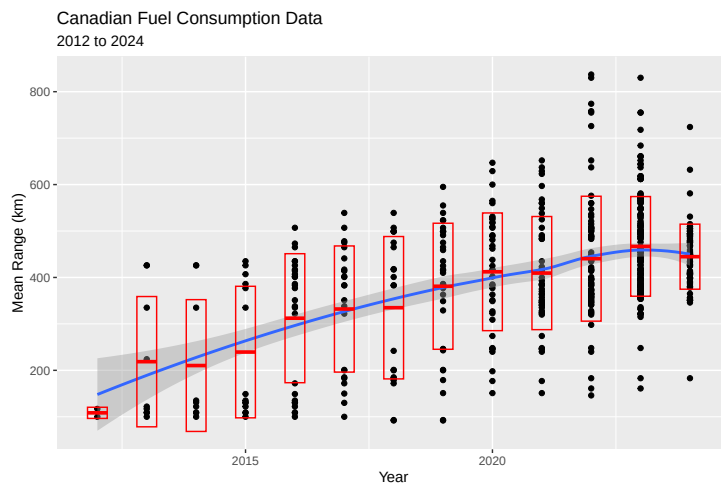


```
...  
stat_summary(  
  fun.data=mean_sdl,  
  fun.args=list(mult=1),  
  color='red',  
  geom="errorbar") +  
...
```



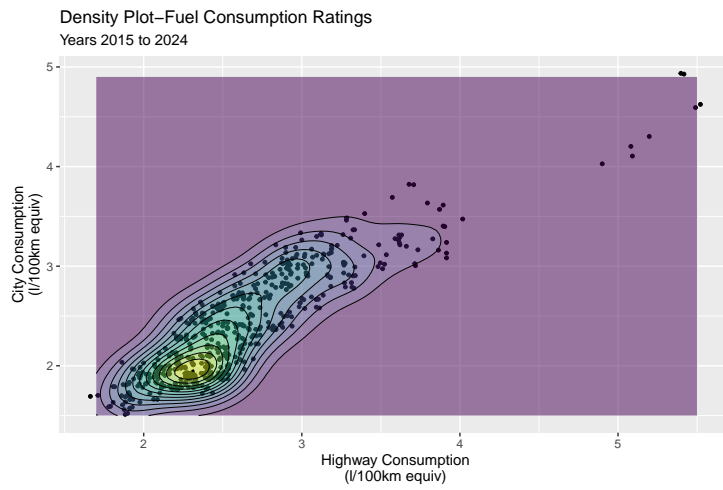
```
...
  stat_summary(
    fun.data=mean_sdl,
    fun.args=list(mult=1),
    color='red',
    geom="crossbar",
    width=0.4) +
...

```



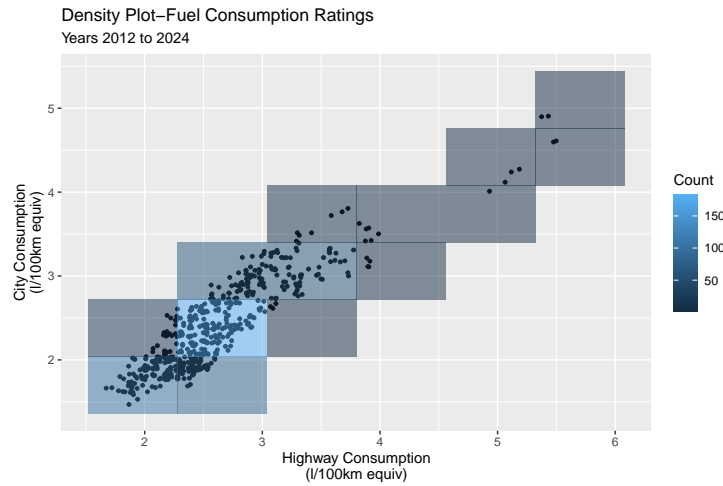
The one-dimensional density plots seen earlier can be generalized to show the two-dimensional joint distribution of values of two variables. The following example uses two geoms for this, one to show the density lines (`geom_density_2d()`), and one to fill the lines in shades of color (`geom_density_2d_filled()`). Additionally, individual data points are plotted using the point geom.

```
e.clean %>%
  ggplot(aes(x=Hwy, y=City)) +
  geom_point(color="black", size=1,
             position='jitter') +
  geom_density_2d_filled(alpha=0.5) +
  geom_density_2d(linewidth=0.25, colour='black') +
  scale_x_continuous(labels=scales::comma) +
  labs(x = 'Highway Consumption\n(l/100km equiv)',
       y = 'City Consumption\n(l/100km equiv)',
       title='Density Plot-Fuel Consumption Ratings',
       subtitle='Years 2015 to 2024') +
  theme(legend.position='none')
```

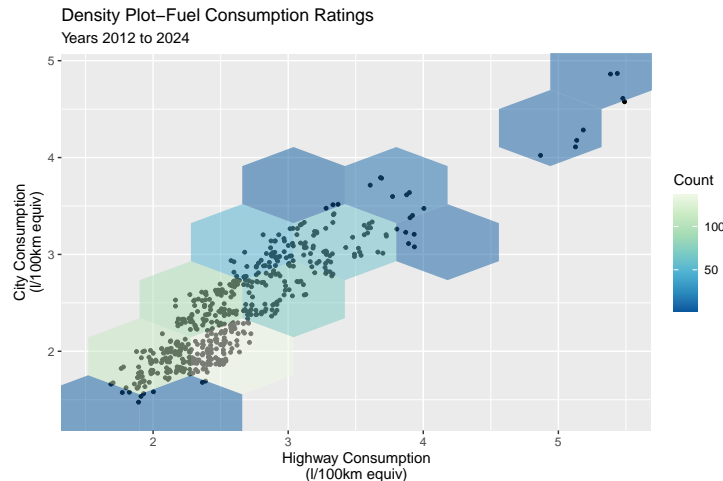


When one is interested in discretizing the two variables, one may show the frequencies or counts in a two-dimensional bin plot. Again, individual data points are shown using the point geom. A somewhat different version is shown below in two-dimensional hex plot, using hexagonal tiles and a different color scale. However, the same information is visualized.

```
e.clean %>%
  ggplot(aes(x=Hwy, y=City)) +
  geom_point(color="black", size=1,
             position='jitter') +
  geom_bin2d(alpha=0.5, bins=5) +
  scale_x_continuous(labels=scales::comma) +
  labs(x = 'Highway Consumption\n(l/100km equiv)',
       y = 'City Consumption\n(l/100km equiv)',
       fill='Count',
       title='Density Plot—Fuel Consumption Ratings',
       subtitle='Years 2012 to 2024')
```

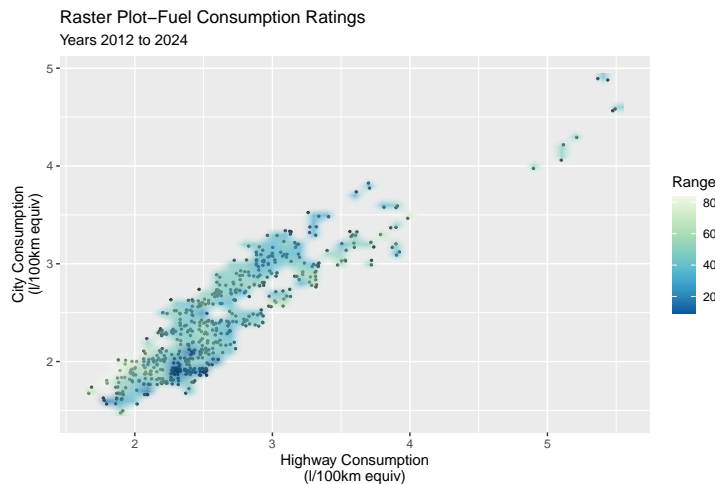


```
...
  geom_hex(alpha=0.5, bins=5) +
  scale_fill_distiller(palette=4, direction=-1) +
...
...
```



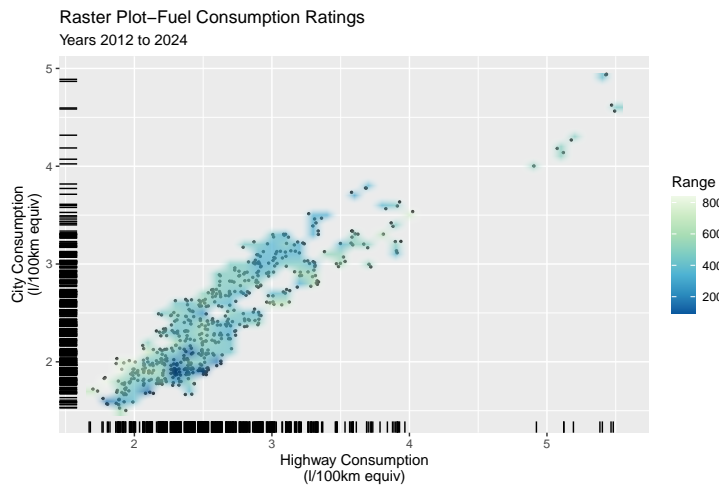
Note that the above two plots are essentially plots of two variables. If, instead of count or density, a third variable is to be shown, one can use the raster geom `geom_raster()`. This requires the aesthetic to specify a data variable mapping for the color element of the plot, as in the following example. Again, individual data points are included with the point geom.

```
e.clean %>%
  ggplot(aes(x=Hwy, y=City)) +
  geom_point(color="black", size=0.5,
             position='jitter') +
  geom_raster(aes(fill=Range), alpha=0.7,
              interpolate=TRUE) +
  scale_fill_distiller(palette=4, direction=-1) +
  scale_x_continuous(labels=scales::comma) +
  labs(x = 'Highway Consumption\n(l/100km equiv)',
       y = 'City Consumption\n(l/100km equiv)',
       fill='Range',
       title='Raster Plot-Fuel Consumption Ratings',
       subtitle='Years 2012 to 2024')
```



The last example shows the use of so-called "rugs" that show marginal distributions of the plot variables. They can be added to different types of plots, including 3D raster plots as done here, as well as 2D bin and hex plots, or one-dimensional histograms. Rugs are added by using the `geom_rug()` function.

```
e.clean %>%
  ggplot(aes(x=Hwy, y=City)) +
  geom_point(color="black", size=0.5,
             position='jitter') +
  geom_raster(aes(fill=Range), alpha=0.7,
              interpolate=TRUE) +
  geom_rug(position='jitter') +
  scale_fill_distiller(palette=4, direction=-1) +
  scale_x_continuous(labels=scales::comma) +
  labs(x = 'Highway Consumption\n(l/100km equiv)',
       y = 'City Consumption\n(l/100km equiv)',
       fill='Range',
       title='Raster Plot-Fuel Consumption Ratings',
       subtitle='Years 2012 to 2024')
```



Hands-On Exercises Using the Pagila database files from the previous chapter on data analysis with R, create the following plots using ggplot2/R. Use the appropriate ggplot2 functions to add informative labels for axes, useful legends to the plots, and use suitable color palettes.

1. A histogram and/or density chart of film length by film category
2. A column chart of the mean rental payments for films by film category
 - Add error bars to this chart
3. A scatter plot of total rental payments by week
 - Add a local regression line to this plot
4. A pie or donut chart of rental counts by film rating

7.9 Visualization in Python using Plotly Express

This section demonstrates visualization in Python using Plotly Express. Plotly Express by default produces web-based, i.e. JavaScript based, interactive plots. On the Python side, the diagram is expressed in more primitive graphical descriptions, serialized in a JSON document, which is sent to the web browser, where the Plotly JavaScript library renders them. Interactivity includes the ability to zoom and pan the plot, and to hover over plot elements to get tooltip overlays, e.g. specific values of points or lines in the plots.

The examples in this section use the same data set as the R examples above, and as much as possible try to provide similar diagrams. The first Python code fragment imports the required packages and loads the data set.

```

import pandas as pd
import plotly.express as px
import plotly.io as pio
pio.kaleido.scope.mathjax = None

# Read data
fuel = pd.read_csv('https://evermann.ca/busi4720/fuel.csv')

```

The `histogram` function does as its name suggests, and produces a histogram. The figure must be shown and can be written to a file using a variety of formats. By default `show()` opens the standard web browser to show the figure. In this mode, the figures are interactive, and can be manually saved to a PNG file.

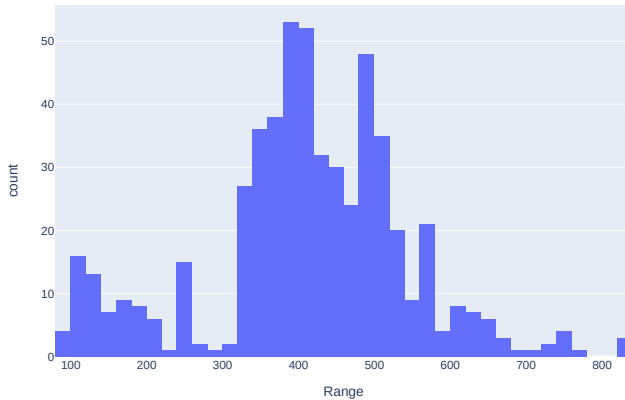
```

# Create histogram
fig = px.histogram(fuel, x='Range', nbins=50)

# Show histogram, by default show in web browser
fig.show()

# Save figure to image
fig.write_image("px.histogram.pdf", height=500, width=750)

```



Similar to the R example earlier, summary information can be added to figures. This is done using the `add_vline()` functions, in the example below with different line styles, annotation text, and annotation positions. The example below also uses `update_layout()` for adding a plot title and providing more informative labels for the x and y axes.

```

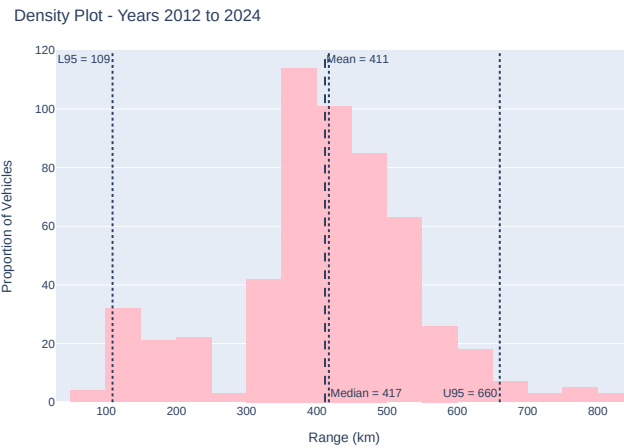
# Calculating summary statistics
mean_v = fuel['Range'].mean()
median_v = fuel['Range'].median()
lower95 = fuel['Range'].quantile(0.025)
upper95 = fuel['Range'].quantile(0.975)

# Creating the density plot
fig = px.histogram(fuel, x='Range',
                    color_discrete_sequence=['pink'])

# Adding vertical lines and annotations
fig.add_vline(x=mean_v, line_dash='dash',
               annotation_text=f'Mean = {round(mean_v)}',
               annotation_position='top right')
fig.add_vline(x=median_v, line_dash='dot',
               annotation_text=f'Median = {round(median_v)}',
               annotation_position='bottom right')
fig.add_vline(x=lower95, line_dash='dot',
               annotation_text=f'L95 = {round(lower95)}',
               annotation_position='top left')
fig.add_vline(x=upper95, line_dash='dot',
               annotation_text=f'U95 = {round(upper95)}',
               annotation_position='bottom left')

fig.update_layout(
    title='Density Plot - Years 2012 to 2024',
    xaxis_title='Range (km)',
    yaxis_title='Proportion of Vehicles')

```



The following column chart example uses the mean fuel consumption information. The `fuel_grouped` DataFrame is prepared by aggregating within groups using the `NamedAgg()` function of Pandas dataframes, forming new columns in the DataFrame for the aggregated information. These columns are then "melted" using the `melt` function so that instead they become rows with names "metric" and "consumption" instead,

where "metric" contains the type of value (city or highway fuel consumption) and "consumption" contains the actual numeric value. The values in the "metric" column are then mapped to new values using the `map()` function on the DataFrame column. This is done so that the chart legend shows "nice" labels. The resulting DataFrame is then used in the `px.bar()` function to produce a bar chart with columns next to each other (`barmode='group'`), informative labels and a title. Note that labels are set in `px.bar` to ensure that tooltip labels on hover are displayed nicely. Finally the figure layout is updated with x and y axes titles.

```

fuel_grouped = fuel.groupby('Year').agg(
    meanCity=pd.NamedAgg('City', 'mean'),
    meanHwy=pd.NamedAgg('Hwy', 'mean')).reset_index()

fuel_long = pd.melt(fuel_grouped,
                    id_vars=['Year'],
                    value_vars=['meanCity', 'meanHwy'],
                    var_name='metric',
                    value_name='consumption')

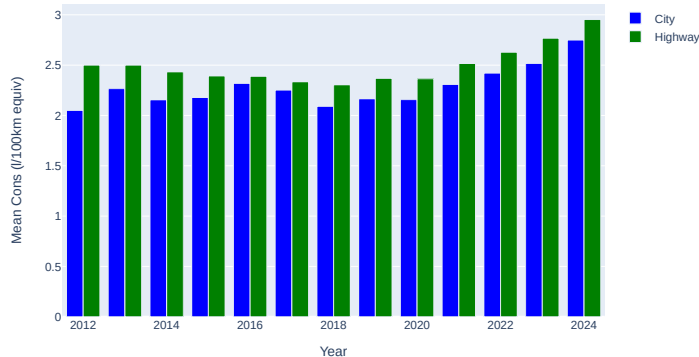
fuel_long['metric'] = fuel_long['metric'] \
    .map({'meanCity': 'City', 'meanHwy': 'Highway'})

fig = px.bar(fuel_long, x='Year', y='consumption',
             color='metric', barmode='group',
             labels={'consumption': 'Mean Cons\n(l/100km equiv)', 'metric': ''},
             title='Electric Vehicle Range (2012 to 2024)',
             color_discrete_map={'City': 'blue', 'Highway': 'green'})

fig.update_layout(
    xaxis_title='Year',
    yaxis_title='Mean Cons\n(l/100km equiv)')

```

Electric Vehicle Range (2012 to 2024)

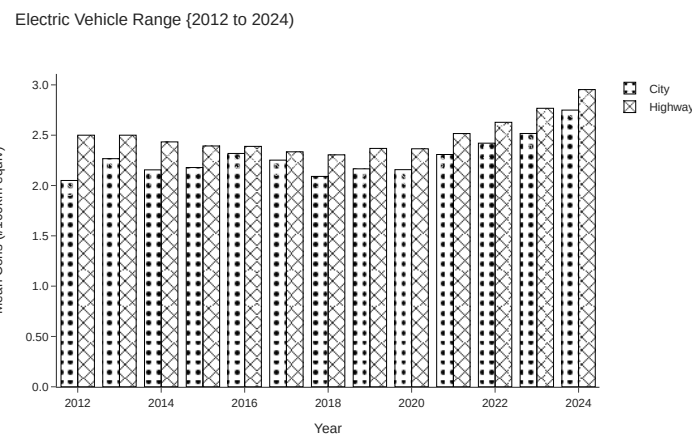


Patterns are easier to do in Plotly Express than in R. Below is the same plot, but using

the `pattern_shape` attribute for the fuel efficiency metric. As for the R example earlier, the sequence of pattern shapes for different categories is provided. Here too, more than the required two are shown to demonstrate the full set of options. The template is set to `simple_white` so as not to show the grid lines or filled background that can be seen in the above plots. The `update_yaxes()` function sets the tick labels to two significant digits, right justified. The `update_traces()` function sets the pattern to black, the fillmode to transparent and

```
fig = px.bar(fuel_long, x='Year', y='consumption',
    pattern_shape = 'metric', barmode='group',
    pattern_shape_sequence = ['.', 'x', '+', '|', '-', '/'],
    title = 'Electric Vehicle Range {2012 to 2024}',
    text_auto=True,
    template="simple_white",
    labels={'consumption': 'Mean Cons\n(1/100km equiv)', 'metric': ''})

fig.update_yaxes(tickformat=',.2r')
fig.update_traces(
    marker=dict(color='black', line_color='black',
                pattern_fillmode='replace'))
```



A box plot is created using the `px.box()` function. Again, the DataFrame is first "melted" from wide format to long format to be able to compare city and highway fuel consumption numbers in the box plot. The example below uses the `update_layout()` function to provide extra information for placing the legend in horizontal format at the top of the plot and centered along the x axis.

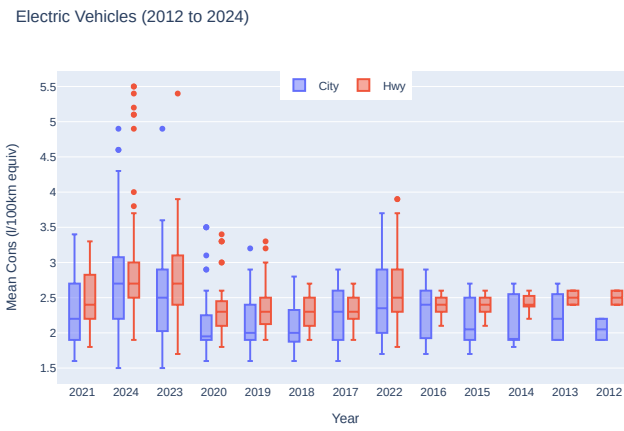
```

fuel_long = pd.melt(fuel,
    id_vars=['Year'], value_vars=['City', 'Hwy'],
    var_name='metric', value_name='consumption')

fig = px.box(fuel_long,
    x=fuel_long['Year'].astype(str),
    y='consumption', color='metric',
    labels={'consumption': 'Mean Cons\n(l/100km)', 'metric': ''},
    title='Electric Vehicles (2012 to 2024)')

fig.update_layout(
    xaxis_title='Year',
    yaxis_title='Mean Cons\n(l/100km equiv)',
    legend_title_text='',
    legend=dict(orientation="h",
                yanchor="top", y=1,
                xanchor="center", x=0.5))

```



A violin plot is constructed similarly to a box plot. Because only the combined fuel consumption numbers are to be shown, there is no variable to be mapped to the "color" plot attribute, and the DataFrame does not need to be transformed first. The `update_traces()` function is used to set the color of the "violins" to black and make them somewhat transparent (`opacity=0.5`). Individual points are shown with a slight jitter to make them distinguishable.

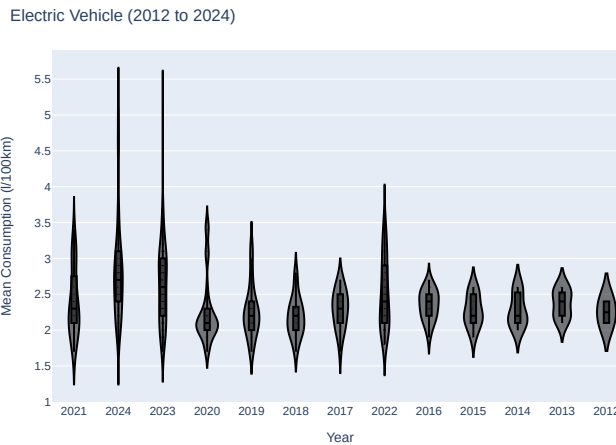
```

fig = px.violin(fuel,
                 x=fuel['Year'].astype(str),
                 y='Comb', box=True,
                 points='all')

fig.update_traces(jitter=0.15, pointpos=0,
                   marker=dict(color='black', size=1, opacity=0.5))

fig.update_layout(xaxis_title='Year',
                  yaxis_title='Mean Consumption\n(l/100km)',
                  title='Electric Vehicle (2012 to 2024)',
                  legend_title_text='')

```



For a count plot, the data frame is first grouped, and the size (that is, the count of values) of each group is recorded in a new column "counts". This transformed data frame is then used for a scatter plot that maps the "counts" variable to the size of the points in the `px.scatter()` function.

```

count_df = fuel.groupby(['Year', 'Category']) \
.size().reset_index(name='counts')

fig = px.scatter(count_df,
                 x='Year', y='Category', size='counts',
                 color_discrete_sequence=['darkolivegreen'],
                 labels={'Category': '', 'Year': 'Year', 'counts': 'Count'},
                 title='EV Models by Category (2012 to 2024)')

```

EV Models by Category (2012 to 2024)

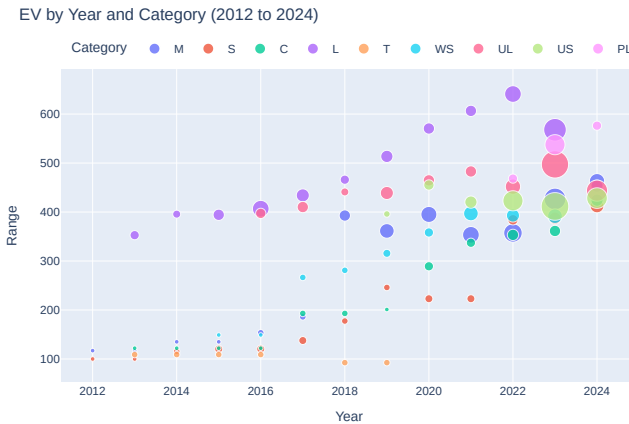


The following points plot is another example of the use of the `px.scatter()` function, which adds a fourth variable to the plot: The vehicle category is mapped to the color plot element. Note also that `update_layout()` is used to provide axis title, create a legend for the "Category" variable (that is mapped to colour), and make the legend horizontal at the top of the plot, right-justified above the x axis.

```
grouped_fuel = fuel.groupby(['Year', 'Category']).agg(
    totalcount=pd.NamedAgg('Range', 'size'),
    meanRange =pd.NamedAgg('Range', 'mean')).reset_index()

fig = px.scatter(grouped_fuel,
                 x='Year', y='meanRange', size='totalcount',
                 color='Category', hover_name='Category',
                 labels={'meanRange': 'Range', 'totalcount': 'Number of Models'},
                 title='EV by Year and Category (2012 to 2024)',
                 size_max=20, opacity=0.8)

fig.update_layout(
    xaxis_title='Year',
    yaxis_title='Range',
    legend_title_text='Category',
    legend=dict(orientation="h", yanchor="bottom",
               y=1.02, xanchor="right", x=1))
```

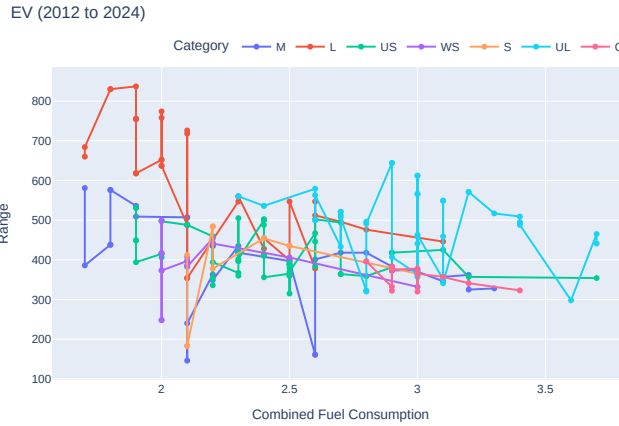


To reduce the amount of information in the following line plot, the data frame is filtered to limit the data set. The `px.line()` function creates the line plot, the option `markers=True` adds the points to the lines. Again, the horizontal legend for the "Category" variable is moved to the top of the plot, right-justified above the x axis.

```
filtered_fuel = \
    fuel[(fuel['Year'] >= 2022) & (fuel['Year'] <= 2023)]
filtered_fuel = filtered_fuel[filtered_fuel['Comb'] <= 4]
filtered_fuel = \
    filtered_fuel[~filtered_fuel['Category'].isin(['PL', 'T'])]

fig = px.line(filtered_fuel,
    x='Comb', y='Range', color='Category',
    line_group='Category', markers=True,
    labels={'Range': 'Range', 'Comb': 'Combined Fuel Consumption'},
    title='EV (2012 to 2024)')

fig.update_layout(
    xaxis_title='Combined Fuel Consumption',
    yaxis_title='Range',
    legend_title_text='Category',
    legend=dict(orientation="h", yanchor="bottom",
               y=1.02, xanchor="right", x=1))
```



Pie charts in Plotly Express are created using the `px.pie()` function. To reduce the number of pie slices to show, the data frame is first filtered for those vehicle makes with 5 or more models and limited to the 2023 model year. The pie chart then shows the number of models for each manufacturer. Labels in the pie chart must be set individually for each pie slice using a `for` loop over the grouped DataFrame rows. For each group (row), an annotation is added to the figure using `add_annotation` that shows the "totalcount" value. Finally, the figure layout is updated to show the legend at the top, right-justified above the x axis.

```

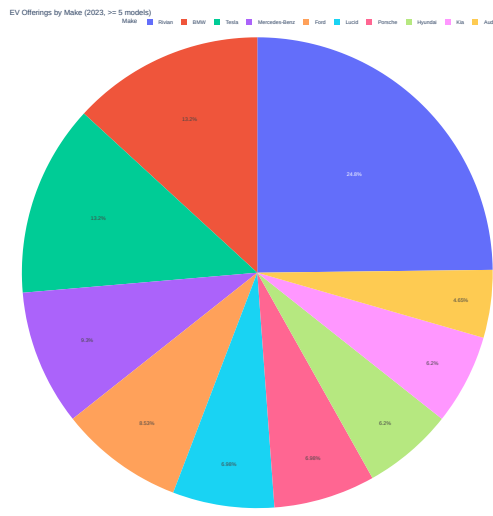
fuel_2023 = fuel[fuel['Year'] == 2023]
fuel_grouped = \
    fuel_2023.groupby('Make').size().reset_index(name='totalcount')
fuel_grouped = fuel_grouped[fuel_grouped['totalcount'] >= 5]

fig = px.pie(fuel_grouped,
             names='Make', values='totalcount', hole=0,
             title='EV Offerings by Make (2023, >= 5 models)',
             labels={'totalcount': 'Number of Models'})

for i, row in fuel_grouped.iterrows():
    fig.add_annotation(text=str(row['totalcount']),
                       x=row['Make'], y=row['totalcount'],
                       showarrow=False, font_color='lightgrey')

fig.update_layout(legend=dict(orientation="h", yanchor="bottom",
                             y=1.02, xanchor="right", x=1),
                  showlegend=True, legend_title_text='Make')

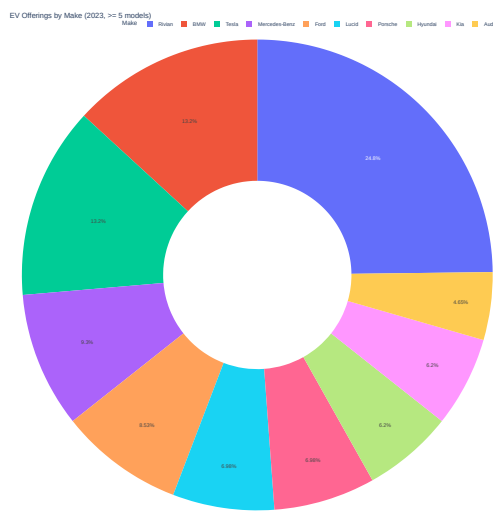
```



A donut chart is easily created simply by providing the `hole` argument to `px.pie()`, as in the following example:

```
...
fig = px.pie(fuel_grouped,
    names='Make', values='totalcount', hole=0.4,
    title='EV Offerings by Make (2023, >= 5 models)',
    labels={'totalcount': 'Number of Models'})
...

```



A radar plot is created as a line plot on a polar coordinate system, using the `px.line_polar()` function. First, the data frame is grouped by vehicle make and aggregate statistics are computed. The grouped data is then filtered to reduce the number of information shown in the radar plot. The aggregate values are then scaled to values between 0 and 1, using the `MinMaxScaler` from the `sklearn` package. Finally, the data frame is "melted" into long format and then used for creating the radar plot shown below.

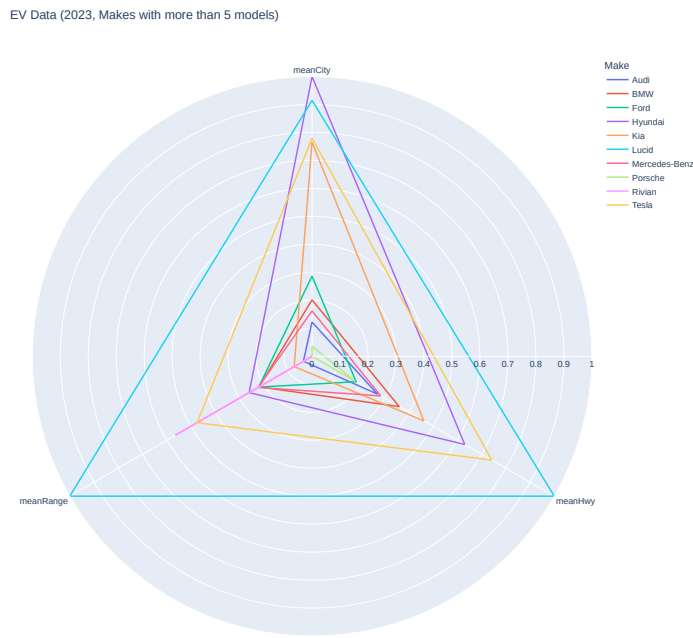
```
from sklearn.preprocessing import MinMaxScaler

fuel_2023 = fuel[fuel['Year'] == 2023]
grouped = fuel_2023.groupby('Make').agg(
    meanCity =pd.NamedAgg('City',lambda x: 1/x.mean()),
    meanHwy =pd.NamedAgg('Hwy',lambda x: 1/x.mean()),
    meanRange=pd.NamedAgg('Range',lambda x: x.mean()/100),
    nModels =pd.NamedAgg('Make','size'))
grouped = grouped[grouped['nModels'] >= 5]

grouped[['meanCity', 'meanHwy', 'meanRange']] = \
    MinMaxScaler().fit_transform(
        grouped[['meanCity', 'meanHwy', 'meanRange']])

melted = grouped.reset_index().melt(
    id_vars='Make', var_name='metric',
    value_vars=['meanCity', 'meanHwy', 'meanRange'])

fig = px.line_polar(melted,
    r='value',
    theta='metric',
    color='Make',
    line_close=True,
    labels={'metric': '', 'value': '', 'Make': 'Make'},
    title='EV Data (Makes with more than 5 models)')
```

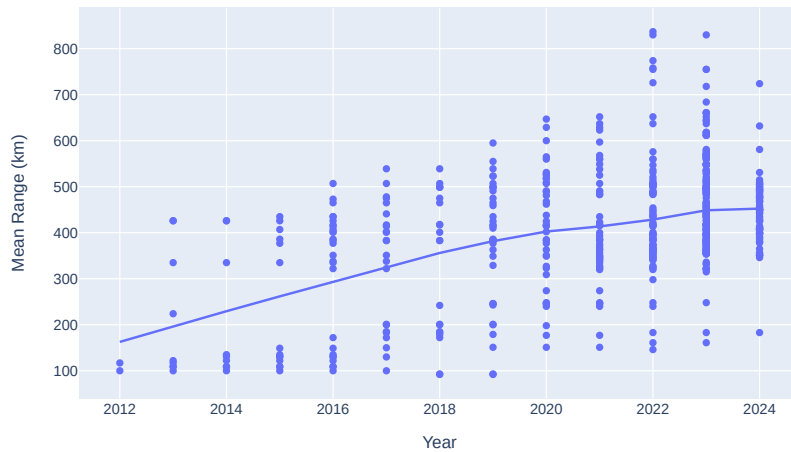


A scatterplot can be created with the `px.scatter()` function which can include trendlines computed using different methods. The most commonly used local regression estimation is specified with the `trendline='lowess'` argument, as in the example below.

```
fig = px.scatter(fuel,
                  x='Year', y='Range', trendline='lowess',
                  labels={'Range': 'Mean Range (km)'},
                  title='EV Range by Year')

fig.update_layout(xaxis_title='Year',
                  yaxis_title='Mean Range (km)')
```

EV Range by Year



Two-dimensional density plots can be created with the `px.density_heatmap()` function, as shown in the following example. This example also uses the `marginal_x` and `marginal_y` options to add histograms for each marginal distribution along the x and y axis. Some other useful options for marginal plots are "rug" and "box".

```
fig = px.density_heatmap(fuel,
    x = 'City', y = 'Hwy',
    nbinsx=20, nbinsy=20,
    color_continuous_scale=px.colors.sequential.Viridis,
    marginal_x="histogram",
    marginal_y="histogram",
    title='EV Fuel Consumption Data',
    labels={"range": "Range",
            "Hwy": "Highway Economy",
            "City": "City Economy"})
```



Hands-On Exercises

Using the Pagila database files from the previous chapters, create the following plots using Plotly Express/Python. Use the appropriate Plotly Express functions to add informative labels for axes, useful legends to the plots, and use suitable color palettes.

1. A histogram and/or density chart of film length by film category
2. A column chart of the mean rental payments for films by film category
 - Add error bars to this chart
3. A scatter plot of total rental payments by week
 - Add a local regression line to this plot
4. A pie or donut chart of rental counts by film rating

7.10 Review Questions

The following review questions are intended to check your understanding of the material on visualization.

1. Explain the significance of data visualization in modern data analysis and communication.
2. How does data visualization blend artistic creativity with analytical skills?
3. List and explain the main reasons why data visualization is used.
4. What is visual discovery in the context of data visualization?
5. Contrast declarative visualization with visual discovery in terms of their purpose and interactivity.
6. Define operational visualization and its role in monitoring and decision making.
7. Explain the importance of focusing on quantitative messages in visualization. Provide examples of how different types of graphs or charts convey different

- types of data or relationships.
8. Discuss some of the challenges or pitfalls that can occur in data visualization, especially regarding pattern recognition and data interpretation.
 9. Explain how the choice of a specific type of data visualization depends on the message or insight that needs to be conveyed.
 10. What are "dark patterns" in the context of data visualization? Provide examples of common dark patterns used to deceive or mislead viewers.
 11. How can cognitive biases be exploited in creating misleading data visualizations?
 12. Explain how scaling and truncating axes in graphs can mislead the viewer. Provide examples.
 13. How can the choice of an inappropriate graph type lead to misleading conclusions? Give specific examples.
 14. Describe how the use of color in data visualization can be misleading. What are the best practices in choosing colors for visualizations?
 15. Discuss the problems associated with using 3D elements or images in graphs. How can these elements distort the data representation?
 16. Describe the unique challenges of visualizing streaming or real-time data. How do these challenges impact the design of such visualizations?
 17. What are the specific challenges of visualizing network or graph data? How do these challenges influence the choice of visualization techniques?
 18. Describe the different types of graph layouts (force-directed, circular, arc, layered) and their use cases. What are the benefits and drawbacks of each layout?
 19. Why are interactive features like zooming, panning, and highlighting important in graph visualizations, especially for large datasets?
 20. List and explain the criteria for assessing the quality of a graph visualization. Why are these criteria important?
 21. Discuss the challenges associated with projecting three-dimensional Earth onto a two-dimensional surface in map visualizations. How do different projections affect the representation of spatial data?
 22. Discuss the techniques used to represent attributes of nodes and edges in network visualizations. How can these techniques enhance or hinder the understanding of the network?
 23. Explain how different areal units (e.g., counties, postal codes, districts) can impact the interpretation of geospatial data visualizations.
 24. Explain why color choice is crucial in data visualizations and list the desirable characteristics of color palettes.
 25. Describe sequential color palettes and discuss their appropriate use cases. Provide an example where a sequential palette is suitable.
 26. What are diverging color palettes and when are they most effectively used in data visualization? Illustrate with an example.
 27. Explain spectral color palettes and their application in visualizing data. Discuss the potential drawbacks of using spectral palettes.
 28. Discuss the importance of considering color vision deficiency (CVD) in choosing color palettes for data visualizations.
 29. How do the different types of color vision deficiencies (e.g., protanopia, deutanopia, tritanopia) affect the perception of colors in data visualizations?

30. Define and discuss the importance of perceptual uniformity in color palettes. How does it impact the interpretation of data?
31. What are monochromatic color palettes and in what situations might they be preferred?
32. What is a box plot and what are the key summary statistics it displays?
33. Explain the concept of the interquartile range (IQR) in a box plot. How is it calculated and what does it represent?
34. Describe the significance of the median line in a box plot. How can the median line's placement provide insights into data skewness?
35. What do the whiskers in a box plot represent? Explain the common method for determining their length.
36. How are outliers represented in a box plot? What criteria is typically used to classify a data point as an outlier in this context?
37. How can you determine if a dataset is symmetric or skewed based on its box plot?
38. Compare and contrast box plots and histograms. In what scenarios might one be preferred over the other?
39. Compare and contrast box plots and violin plots. In what scenarios might one be preferred over the other?

Chapter 8

Business Process Analytics

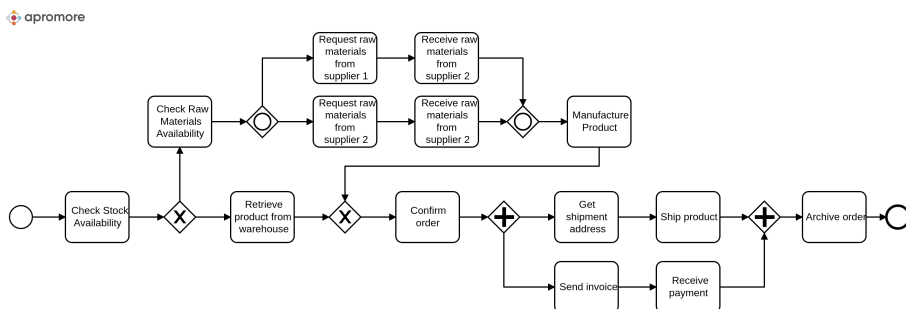
8.1 Introduction

Business process analytics is concerned with using event data to improve the *operational efficiencies* of business processes. A business process is how an organization creates value for its customers. Improvements to operations may yield better customer service and increase customer satisfaction, it may also reduce the time it takes to complete a business process, it may reduce employee workload and improve employee satisfaction, it may reduce cost, or free up capacity.

Every organization manages and executes a multitude of business processes. An important business process in many organizations is the order-to-cash process, which is the sequence of activities that begins when a customer submits an order and ends when the money has been received and a receipt has been issued. Another important process is issue-to-resolution, which begins when a customer contacts the organization with a problem about a product or service, and ends when the problem has been resolved to the satisfaction of the customer. Business processes exist in all kinds of organizations, from for-profit manufacturing or service enterprises, to healthcare clinics and hospital, to education and government services, to non-profit and charity organizations. Typically, a business process is characterized by its *business object*. For example, the business object for the order-to-cash process is the customer order, the business object for the issue-to-resolution process is the customer complaint.

8.2 Business Processes and Business Process Models

A business process is defined as a *sequence of activities* that are executed in a defined order to create some type of *value for a customer*. Besides activities, a business process also includes the *resources* that carry out activities. Resources may be human resources, that is people or employees, or they may be machines or computer systems.



Source: Marlon Dumas, Marcello La Rosa, Jan Mendling, Hajo A. Reijers (2018) "Fundamentals of Business Process Management", 2nd edition, Springer Verlag. Figure 3.12.

Figure 8.1: Example BPMN model

Resources typically play one or more *roles* in organization, such as accountants, warehouse workers, service technician, etc.

A business process may also contain *events*. Events can trigger or start a business process or they may occur within a business process. Typical events are customer orders arriving, customer inquiries arriving, goods or materials arriving, goods or materials being dispatched, etc.

A business process may also contain *decisions*. For example, in an order-to-cash process, a decision may need to be made how to source a part, or how to ship a part, or whether to invoice a customer, etc. In a healthcare process, decisions may involve treatment options, medical tests, or hospital admissions.

Business processes are typically defined using business process models or business process diagrams. A common way to describe processes is with the *Business Process Modelling Notation (BPMN)*, the industry standard developed by the OMG (Object Management Group)¹ and adopted as the ISO/IEC 19510 standard. Figure 8.1 shows an example of a business process model in BPMN.

In this model, the circles represent events, rectangles represent activities, diamond shapes represent "gateways" and arrows represent process flow *dependencies*. Dependencies describe what must happen before something else can happen, or, alternatively, what can follow once something is completed.

The diamond with the "×" symbol represents an *exclusive gateway*, which means that a decision is made and only one outgoing path can be taken by the process. For example, after the activity "Check Stock Availability", either the activity "Check Raw Materials Availability" is carried out, or the "Retrieve product from warehouse" activity.

In contrast, the "+" symbol represents a *parallel gateway*, which means that the process proceeds along *all* outgoing paths, in any order, or possibly at the same time. For

¹<https://www.omg.org/bpmn/>

example, after the "Confirm order" activity, both the "Get shipment address" and the "Send invoice" activities are carried out, in any order and possibly at the same time.

Finally, the "○" symbol represents an *inclusive gateway*. The process may proceed along any number of outgoing paths. For example, after the "Check Raw Materials Availability" activity, the process may proceed with the "Request raw materials from supplier 1" activity, or the "Request raw materials from supplier 2" activity, or both of them.

8.3 Business Process Event Logs

A *case* is one instance of a business process. For example, the execution of the order-to-cash process and its activities for the order number 1234 is one case (instance); executing the process for order number 2345 is another case (instance).

A *trace* is the sequence of events for one case. An *event* in this context is usually the execution of an activity instance (for example, "Check Stock Availability" for order 1234) or the occurrence of an outside event (for example, "goods have arrived at warehouse"). However, activity instances themselves can have a complicated lifecycle. Figure 8.2 shows an example of the lifecycle model of the XES standard² for event log data. The XES standard defines how event data is represented in an XML document.

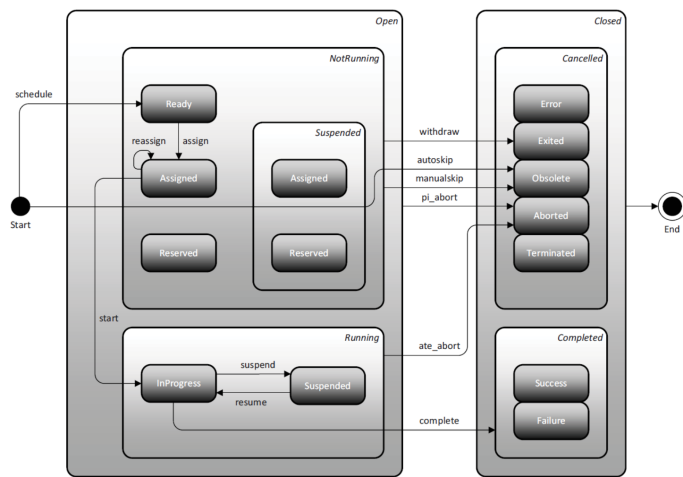
Each arrow in this figure represents a lifecycle transition and each box represents a lifecycle state. For example, an activity instance is first scheduled, then assigned to a resource, then started by the resources, and finally completed successfully. However, other lifecycles are possible in the XES lifecycle model. For example a resource may be repeatedly reassigned before being started, it may be suspended and resumed, and it may be skipped or aborted. Each of these lifecycle transitions may be captured by an *event* in an event log.

Each event may be associated with additional information, either about the particular activity instance or about the case. These are called *event attributes* or *case attributes*. Case attributes in an order-to-cash process instance may be name of the customer, the list of ordered products, etc. An example of an event attribute for a "Confirm order" activity instance completion event is the confirmation number that is generated by the activity instance. Each event is also typically associated with information about the resource that executed the activity instance that the event refers to, as well as a timestamp of when the event occurred.

An *event log* is a collection of one or more traces for one process. Hence, an event log describes the execution of one or more cases of the same process. Note that event logs may contain incomplete cases (e.g. cases that have not been completed when the event log was collected), may be randomly sampled from the complete event log data, etc.

Event logs are typically generated by process-aware information systems such as dedicated workflow-management systems, but many other corporate information systems,

²<https://www.tfpm.org/resources/xes-standard>



Source: <https://www.tf-pm.org/resources/xes-standard/about-xes/standard-extensions/lifecycle/standard>

Figure 8.2: Process Activity Lifecycle

such as Enterprise Resource Planning (ERP) systems, Supply Chain Management (SCM) systems, Customer Relationship Management (CRM) systems, and other, keep track of *who did what when*, which is the basic information in any event log. Additionally, event logs may be collected from any web-based information system as web-servers routinely keep log information about user interaction with the web site.

To be usable for process analytics, the event log information from source information systems must typically be extracted, transformed and then loaded into a format that is used by process analytics software. This is known as an *ETL process*: Extraction–Transformation–Load. ETL processes are required for many analytics applications that take their raw data from a variety of sources.

The standardized interchange format for event log data is the XES file format. XES stands for eXtensible Event Stream and is an XML based format. Another common format used for event log data are CSV files, where each row typically corresponds to one activity instance (not one event!). Below is an example of an XES file that uses the standard XES activity lifecycle model, defines three case attributes ("REG_DATE", "AMOUNT_REQ", and "concept:name") and three event attributes ("time:timestamp", "lifecycle::transition" and "concept:name" and then contains traces with their events. The XML code below is an excerpt of an XES file and illustrates how traces and events are encoded in XML.

```

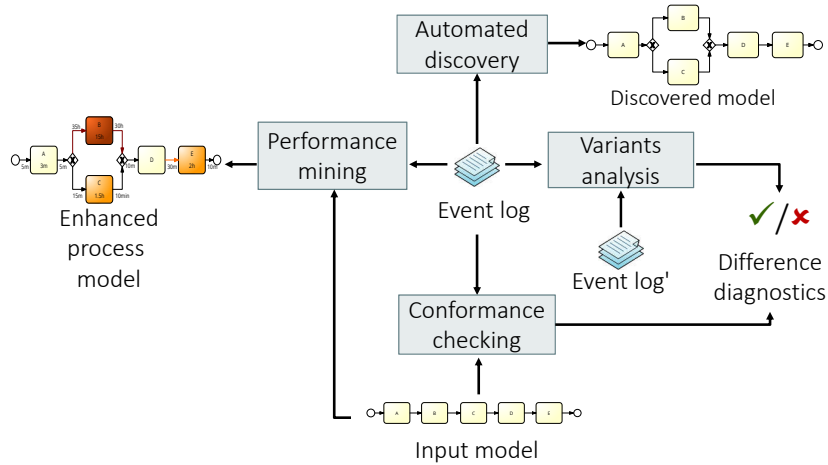
<?xml version="1.0" encoding="UTF-8" ?>
<log xes.version="1.0" xes.features="nested-attributes"
      openxes.version="1.0RC7"
      xmlns="http://www.xes-standard.org/">
  <extension name="Lifecycle" prefix="lifecycle">
```

```

    uri="http://www.xes-standard.org/lifecycle.xesext"/>
<extension name="Organizational" prefix="org"
    uri="http://www.xes-standard.org/org.xesext"/>
<extension name="Time" prefix="time"
    uri="http://www.xes-standard.org/time.xesext"/>
<extension name="Concept" prefix="concept"
    uri="http://www.xes-standard.org/concept.xesext"/>
<global scope="trace">
    <date key="REG_DATE" value="1970-01-01T00:00:00.000+01:00"/>
    <string key="AMOUNT_REQ" value="UNKNOWN"/>
    <string key="concept:name" value="UNKNOWN"/>
</global>
<global scope="event">
    <date key="time:timestamp" value="1970-01-01T00:00:00.000+01:00"/>
    <string key="lifecycle:transition" value="UNKNOWN"/>
    <string key="concept:name" value="UNKNOWN"/>
</global>
<classifier name="Activity classifier"
    keys="concept:name lifecycle:transition"/>
<classifier name="Resource classifier"
    keys="org:resource"/>
<trace>
    <date key="REG_DATE" value="2011-10-01T09:45:37.274+02:00"/>
    <string key="concept:name" value="173706"/>
    <string key="AMOUNT_REQ" value="18000"/>
    <event>
        <string key="org:resource" value="112"/>
        <string key="lifecycle:transition" value="COMPLETE"/>
        <string key="concept:name" value="A_SUBMITTED"/>
        <date key="time:timestamp" value="2011-10-01T09:45:37.274+02:00"/>
    </event>
    <event>
        <string key="org:resource" value="112"/>
        <string key="lifecycle:transition" value="COMPLETE"/>
        <string key="concept:name" value="A_PARTLYSUBMITTED"/>
        <date key="time:timestamp" value="2011-10-01T09:45:37.363+02:00"/>
    </event>
    ...
</trace>
...
</log>
```

The file below is an excerpt of a CSV file that contains event log information (line breaks have been added to fit this into the width of the page but are not in the actual file). Note how "Start Timestamp" and "Complete Timestamp" exist for each row. This means that one row captures two events, the starting and the completion of an activity instance.

Case ID	Start Timestamp	Complete Timestamp	Activity	Resource	Role
339	2011/02/16 14:31:00.000	2011/02/16 15:23:00.000	Create Purchase Requisition	Nico Ojenbeer	Requester
339	2011/02/17 09:34:00.000	2011/02/17 09:40:00.000	Analyze Purchase Requisition	Maris Freeman	Requester Manager



Source: Marlon Dumas, Marcello La Rosa, Jan Mendling, Hajo A. Reijers (2018) "Fundamentals of Business Process Management", 2nd edition, Springer Verlag. Figure 11.3

Figure 8.3: Overview of Process Analytics with Event Logs

```

339,2011/02/17 21:29:00.000,2011/02/17 21:52:00.000,
Amend Purchase Requisition,Elvira Lores,Requester
339,2011/02/18 17:24:00.000,2011/02/18 17:30:00.000,
Analyze Purchase Requisition,Heinz Gutschmidt,Requester Manager

```

8.4 Types and Goals of Process Analytics

The diagram in Figure 8.3 shows four important activities in process analytics. Automated process discovery discovers a process model from an event log. This is useful to understand how a process is actually executed. The discovered process can then be analyzed for weaknesses or compared to a normative process model, e.g. as part of an audit. Many organizations also do not have well-defined processes, so that automated discovery is an important first step in understanding their own operations in detail.

Conformance checking compares an event log to a given input process model. It checks whether the actual operations, as captured in the event log, conform to or comply with a normative process model, a set of business rules, or a set of process constraints. This is typically done as part of an audit to demonstrate compliance, for example in the financial services industry, healthcare, or for quality management certifications.

Performance mining enhances a process model (which could be an automatically discovered one) with information about the duration of activities and waiting times between activities, as captured in an event log. From there, process analysts can identify bottlenecks in the process, that is, activities where cases have to wait to be processed,

activities that take a long time, activities with high variability in their processing time, or similar process problems. Further information can then be collected to identify the causes of and remedies for these problems.

Finally, variants analysis compares two different event logs of the same or similar processes to understand differences in execution. This may be useful when organizations execute the same business processes in different locations, or in different business units. This could be used to identify best practices for an organization.

In addition to these four, process prediction has become an important aspect of business process analytics. Process prediction uses an event log to train a statistical model in order to predict the future course or outcome of a currently running case. Typical prediction targets are the remaining time to completion of the case, the most likely next activity, the probability of a negative or positive outcome, the waiting time before the next activity starts, etc. Process prediction is useful to allow process managers to proactively intervene in a case before a problem arises, or before a case becomes late or overdue. Process prediction can also serve to provide information to customers about the how their case will likely be handled or completed.

Overall, the purposes of process analytics are multiple:

- Discover actual operations
- Check actual process against desired process
- Identify operational (performance) problems
- Improve operational processes
- External compliance analysis and reporting
- Identify implicit or de-facto organizational groups and relationships
- Support, reinforce, or break organizational relationships

8.5 Process Analytics Tools

Since the inception of the field of process analytics circa early 2000s, a range of commercial and open-source tools have been developed to support process analysts. Among the widely-used commercial tools are Celonis, Signavio, Fluxicon, ARIS and Apromore.

Celonis Celonis³ is a leading process mining software that excels in helping organizations analyze and optimize their business processes through powerful data visualization and analytics. This tool extracts and leverages data from various IT systems to provide real-time insights into process performance, identify bottlenecks, and uncover inefficiencies. Celonis facilitates extensive process discovery, conformance checking,

³<https://www.celonis.com/>

and predictive modeling, empowering users to drive significant improvements in process efficiency and effectiveness. Its capabilities are enhanced by features like machine learning and automation, making Celonis a pivotal tool for enterprises aiming to execute large-scale digital transformation strategies and achieve operational excellence.

Signavio Process Manager Signavio⁴ Process Miner, part of the Signavio Business Transformation Suite, enables organizations to analyze and optimize business processes by visualizing actual workflows and identifying deviations from ideal models. This tool automatically generates process models from data logs, performs conformance checks to ensure regulatory compliance, and analyzes process performance metrics such as duration, frequency, and costs. Its integration with Signavio's broader suite allows for a seamless workflow from process discovery through modeling to execution, making it a valuable tool for continuous process improvement and alignment across organizational departments.

Fluxicon Disco Fluxicon Disco⁵ is a user-friendly process mining software that excels in providing fast and intuitive insights into business processes. It is designed for ease-of-use, allowing users to quickly load data and start analyzing with minimal setup. Disco supports a range of features including automated process discovery, performance analysis, and bottleneck identification, making it ideal for users seeking immediate and actionable insights. With its strong focus on visual analytics, Fluxicon Disco offers detailed, interactive process maps and a variety of filters to explore process variations and issues efficiently. This tool is popular among both academic researchers and industry professionals for its simplicity and powerful analytical capabilities.

ARIS ARIS Process Mining⁶ is a robust tool designed to help organizations discover, measure, and analyze their business processes in order to identify inefficiencies and optimize performance. It enables users to visualize complex process flows and pinpoint deviations, bottlenecks, and vulnerabilities by extracting data from IT systems and reconstructing the actual processes that take place. ARIS offers comprehensive analytics capabilities, including conformance checking, root cause analysis, and simulation for predicting process behavior and outcomes. This integration with digital transformation initiatives makes ARIS an important tool for businesses aiming to achieve operational excellence and continuous improvement in their processes.

Apromore Apromore⁷ is a leading cloud-based process mining tool known for its sophisticated analytics capabilities and user-friendly interface. It provides advanced process mining techniques such as automated process discovery, conformance checking, and predictive analytics. Apromore is designed to handle large and complex datasets efficiently, offering deep insights into business processes to help organizations identify

⁴<https://www.signavio.com/>

⁵<https://www.fluxicon.com/>

⁶<https://aris.com/process-mining/>

⁷<https://apromore.com/>

inefficiencies, ensure compliance, and enhance operational performance. Its collaborative features support multi-user environments, making it a good choice for enterprises aiming to undertake continuous process improvement and drive operational excellence through detailed data-driven insights.

Among the widely-used open-source tools are ProM (a system for research) and BupaR (for R) and PM4PY (for Python).

ProM ProM⁸ is a versatile open-source process mining tool that stands out for its extensive range of plugins supporting a diverse array of process mining tasks, including discovery, analysis, and enhancement of business processes. Developed primarily for academic and research purposes, ProM offers functionalities for detailed process discovery, conformance checking, and social network analysis among others. It is highly regarded for its flexibility, allowing researchers and professionals to experiment with new algorithms and techniques through its modular and extensible architecture. ProM's ability to handle various types of event logs and its rich collection of tools make it a good choice for in-depth process mining investigations and experiments.

bupaR bupaR⁹ is an R-based open-source library specifically designed for process mining and business process analysis. It offers a comprehensive suite of tools that enable users to perform detailed process discovery, conformance checking, and performance analysis directly within the R programming environment. bupaR leverages the extensive data manipulation and visualization capabilities of R, allowing users to integrate process analysis with statistical and predictive analytics seamlessly. This makes it a good choice for statisticians and data scientists looking to conduct in-depth process analysis, create interactive process visualizations, and derive actionable insights from process data, all within the familiar and powerful R ecosystem.

PM4Py PM4Py¹⁰ is a Python library that offers a comprehensive suite of process mining tools, making it a powerful resource for performing process discovery, conformance checking, and process enhancement. Tailored for the Python ecosystem, PM4Py facilitates the analysis of complex process data by integrating seamlessly with popular data science tools such as pandas and numpy. Its capabilities extend to generating process models from event logs, analyzing process performance, and providing insights into workflow efficiencies and bottlenecks. Ideal for both academic research and practical applications, PM4Py is known in the process mining community for its accessibility, scalability, and the ease with which users can implement and customize process mining algorithms.

⁸<https://github.com/promworkbench>

⁹<https://bupar.net/>

¹⁰<https://processintelligence.solutions/pm4py>

8.6 Process Mining in Python with PM4Py

This section illustrates process analytics using the PM4Py framework for Python. In a first step, we import an event log in CSV format into a Pandas data frame¹¹. The event log is a fictitious log of a purchasing or procurement process. There are no case or event attributes other than the basic information about activity names, resources, and timestamps.

```
import pandas as pd
import pm4py

# Load the event log and parse date columns
log = pd.read_csv('https://evermann.ca/busi4720/PurchasingExample.csv',
                   parse_dates=['Start Timestamp', 'Complete Timestamp'],
                   infer_datetime_format=True)
```

In order for any process analytics tool to work with event log data, it must know which column in the data set is the case identifier, so that it knows which events belong to each case. It must also know what the name of the activity of each event is, the correct timestamp for sequentially ordering events within a case, and the resource that executed the activity referred to by the event. PM4Py by default uses attribute names similar to those in the XES file above, although others can be explicitly specified. The following Python code block defines the expected columns in the data set with the appropriate names and types.

```
# Tell PM4PY about which columns represent case ID, activity name,
# and timestamp. Case ID and activity. Names must be string type
log['case:concept:name']=log['Case ID'].astype('string')
log['concept:name']=log['Activity'].astype('string')
log['time:timestamp']=log['Complete Timestamp']
log['org:resource']=log['Resource']
```

Reading an XES file into PM4Py is also easy, but because it does not use the Pandas library, XES files must be on a local filesystem (although they may be compressed). As illustrated above, an XES file contains sufficient meta-data to identify case ID, event name, timestamp, and resource, so that these need not be specified upon import.

```
log2 = pm4py.read_xes('BPI_Challenge_2012.xes.gz')
```

Basic Log Information

Basic event log statistics can of course be computed through Pandas data frame operations as in the first two lines of the following example code block, but PM4Py provides

¹¹The event log is originally taken from here: <http://files.fluxicon.com//Datasets/Purchasing-Example.csv>

easy-to-use functions. The following Python code block shows the number of cases, number of events, the set of all start activities of traces, the set of all end activity of traces, case durations, and trace and event attributes. The last two are only applicable to event logs in XES format.

```
num_cases = len(log['Case ID'].unique())
num_events = log.shape[0]

pm4py.get_start_activities(log)
pm4py.get_end_activities(log)

pm4py.get_all_case_durations(log)

# Useful only or XES-based event logs
pm4py.get_event_attributes(log)
pm4py.get_trace_attributes(log)
```

Variants

To perform a variant analysis, the log must be separated into sets of traces ("sub-logs") for which every trace contains the same sequence of events. The PM4Py function `split_by_process_variant()` returns an iterator over the variants and their associated sub-logs, where each variant is the list of activity names in the order in which they were executed in that variant. Each sub-log can then be analyzed separately, for example in compliance analysis or performance mining, allowing deeper insights and also a comparative analysis of processes.

```
pm4py.get_variants(log)

# Split the log into sub-logs
for variant, subdf in pm4py.split_by_process_variant(log):
    print(variant)
    print(subdf)
```

Process Discovery

The basic output of process discovery is a dependency graph, also called directly-follows graph or process map. This graph simply shows how often one activity is followed by another activity in the traces of the log. The PM4Py function `discovery_dfg()` returns the directly-follows-graph and also the set of start and end events. This graph can then be visualized using the `view_dfg()` function, as shown in Figure 8.4, and saved to a file using the `save_vis_dfg()` function. This is illustrated in the following Python code lblock.

```

dfg, start, end = pm4py.discover_dfg(log)

pm4py.view_dfg(dfg, start, end, rankdir='LR')

pm4py.save_vis_dfg(dfg=dfg,
    start_activities=start, end_activities=end,
    file_path='dfg.png', rankdir='TB')

```

To show the usefulness of even this basic process diagram, consider the following observations. First, the example DFG in Figure 8.4 shows that all 608 traces in the log start with the same activity ("Create Purchase Requisition"). The activity "Analyze Request for Quotation" is carried out 1107 times, suggesting that some cases require this activity multiple times, as is also evident from the loops between this activity and the activities "Amend Request for Quotation Requester" and "Amend Request for Quotation Requester Manager". This iteration suggests re-work to fix errors in the original quotation. Eliminating these errors may improve the overall process performance.

Second, the DFG shows that 131 cases end after the request for quotation is analyzed, suggesting that these requests are not approved for purchasing. This implies wasted effort and a process analyst may wish to identify ways to reduce these unsuccessful purchase requisitions.

Third, note that there are 10 cases where the activity "Release Supplier's Invoice" is skipped. This may indicate a potential compliance problem and a process analyst may wish to identify which cases skipped this activity and why they did so, or were allowed to do so.

A DFG is not a process model in the sense of the example BPMN model in Figure 8.1, as it is missing gateways and decisions. A range of algorithms have been developed over the years to discover BPMN models from event logs. PM4Py provides the Inductive Miner and the Heuristics Net Miner.

The Inductive Miner works by repeatedly "cutting" the DFG for an event log to identify subsets of activities that represent exclusive choice, parallelism, sequence, or loops. For example, as Figure 8.5 shows, when there are sets of activities that are not connected by each other, and have distinct pre- and post-sets of activities, then this may indicate an exclusive choice between these sets. Similarly, a sequence of activities is characterized by the absence of "backwards" connections in the DFG, and parallelism is indicated by a set of arrows that fully connects two sets of activities with a common post-set.

PM4Py provides the function `discover_bpmn_inductive` which produces a BPMN model that can be viewed and saved. An example is shown in Figure 8.6.

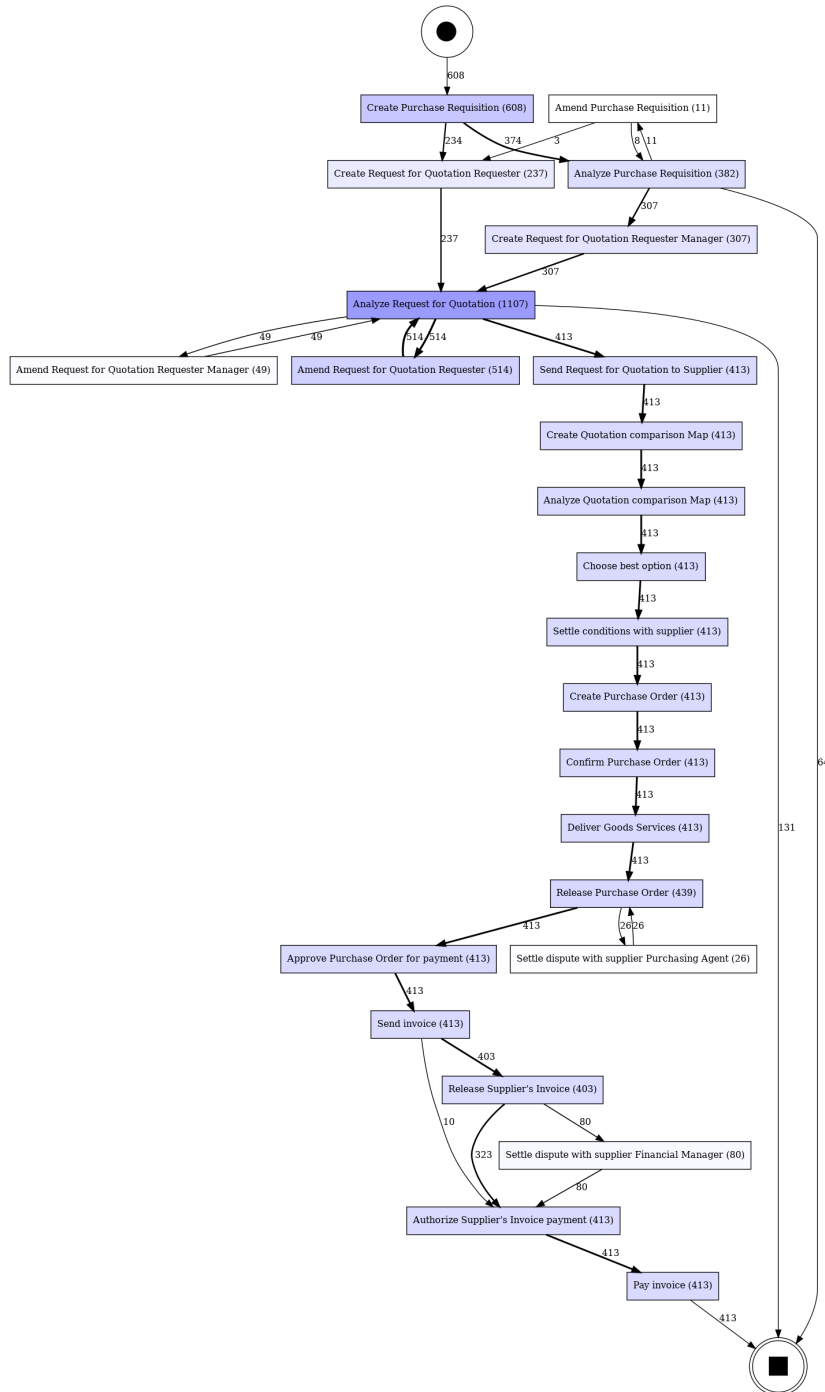
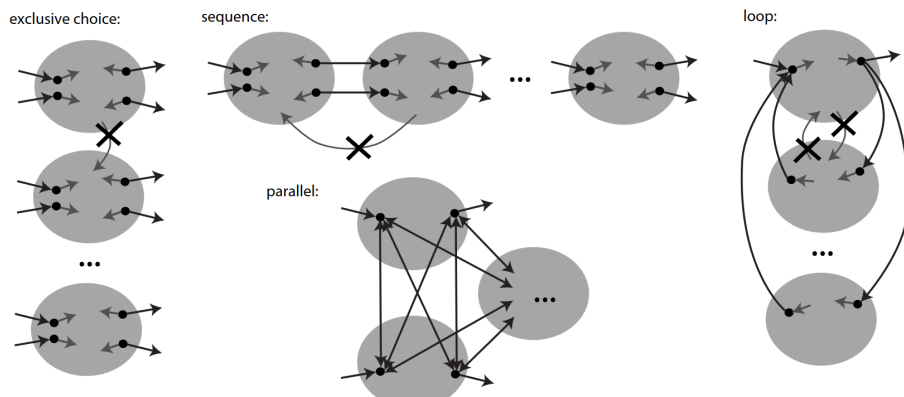


Figure 8.4: Directly-Follows-Graph



Source: Leemans, S.J.J., Fahland, D., van der Aalst, W.M.P. (2013). Discovering Block-Structured Process Models from Event Logs - A Constructive Approach. In: Colom, JM., Desel, J. (eds) Application and Theory of Petri Nets and Concurrency. PETRI NETS 2013. Lecture Notes in Computer Science, vol 7927. Springer, Berlin, Heidelberg.

Figure 8.5: Principles of the Inductive Miner

```
bpmn_model = pm4py.discover_bpnm_inductive(log, noise_threshold=0.5)
pm4py.view_bpnm(bpnn_model, rankdir='LR')
pm4py.save_vis_bpnm(bpnn_model, file_path='bpnn.png', rankdir='TB')
```

In contrast, the Heuristics Net Miner focuses on frequencies in the DFG to identify sequences, parallelism, and loops. For example, a sequence between activities "a" and "b" by values larger than a threshold for the following proportion where the relation $a > b$ indicates the number of times b follows a in the DFG. Similar expressions exist to identify parallelism and loops. The thresholds are configurable; lower thresholds lead to the inclusion of more detail in the final model.

$$a \Rightarrow b = \left(\frac{|a > b| - |b > a|}{|a > b| + |b > a| + 1} \right)$$

The PM4Py function `discover_petrinet_heuristics` provides the Heuristics Net Miner. It returns a Petri net (a type of process model) that can be converted to a BPMN model for viewing and saving. An example is shown in Figure 8.7.

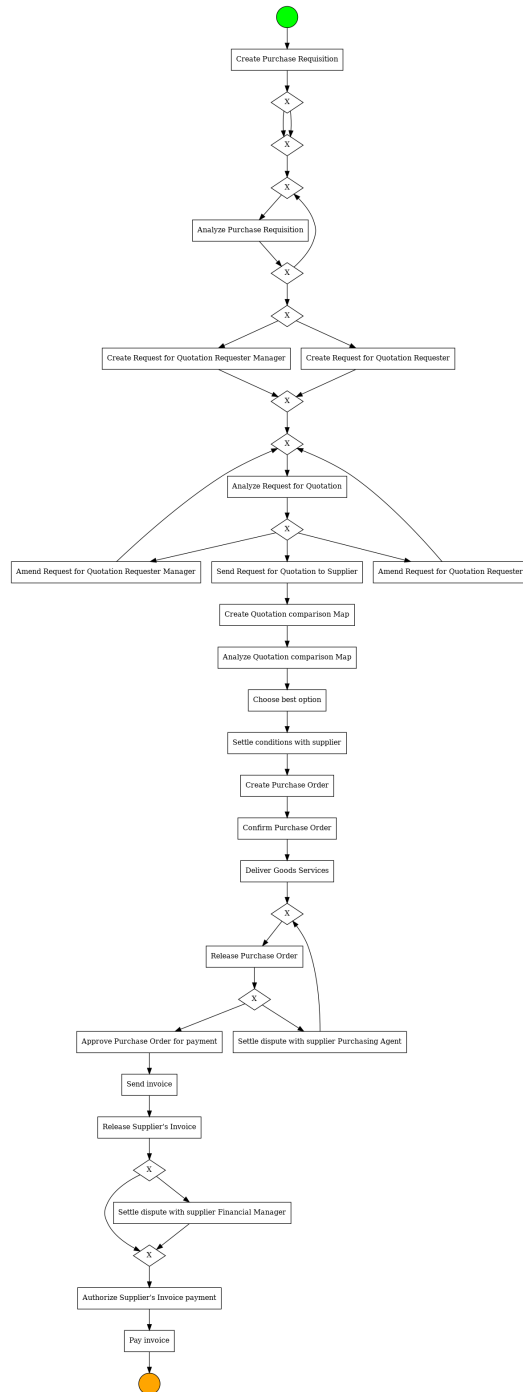


Figure 8.6: Model discovered by the Inductive Miner

```

petri_net, initial_marking, final_marking = \
    pm4py.discover_petri_net_heuristics(log,
        dependency_threshold=0.6,
        and_threshold=0.65,
        loop_two_threshold=0.4)

pm4py.view_petri_net(petri_net)

bpmn_model2 = pm4py.convert_to_bpmn(petri_net,
    initial_marking, final_marking)

pm4py.view_bpmn(bpmn_model2)
pm4py.save_vis_bpmn(bpmn_model2, 'bpmn2.png', rankdir='TB')

```

Once a model is automatically discovered from a log, the analyst must assess its quality. There are four aspects of model quality with respect to an event log:

- **Fitness:** Can the model generate all traces in log?
- **Precision:** Does the model only generate traces in log?
- **Generalization:** Can the model generalize to "sensible" traces not seen in log?
- **Complexity:** Is the model too complex to understand?

Quality assessment often focuses mainly on the calculation of fitness and precision, and two different techniques have been developed for this.

In *token-based replay*, each trace of an event log is replayed on the discovered process model using what is known as *token semantics*. This discovers missing and surplus tokens, which represent model activities that cannot be executed, or model activities that are executed too often. Both cases indicate a mismatch between the model and the trace. The statistics of interest are the percentage of traces that fit the model perfectly, and the average fitness of all traces in the log.

Alignment-based fitness uses sequence alignment methods to align the model and each trace in an event log. It counts the the number of "synchronous moves" where an activity is both in a trace and the model, the number of "move on log" where an activity is in a trace but not in the model, and the number of "move on model" where an activity is in the model but not in the trace. The statistics of interest are also the percentage of traces that fit the model perfectly, and the average fitness of all traces.

Note that the results of the two types of analysis are not necessarily the same for all models and for all event logs.

PM4Py provides functions to compute fitness and precision using both methods. For this, PM4Py uses the Petri net type of process model. As we noted above, this can be transformed into a BPMN model for visualization.

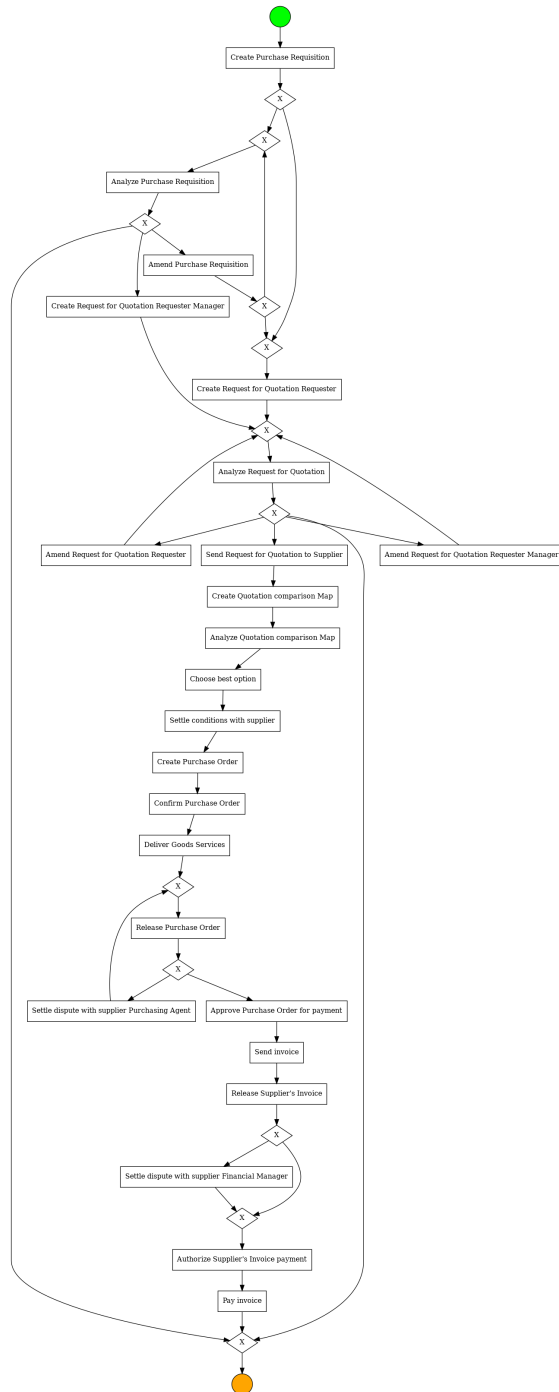


Figure 8.7: Model discovered by the Heuristics Net Miner

```
petri_net, initial_marking, final_marking = \
    pm4py.discover_petri_net_inductive(log, noise_threshold=0.5)

fitness_alignments = pm4py.fitness_alignments(log,
    petri_net, initial_marking, final_marking)
print(fitness_alignments)

fitness_tbr = pm4py.fitness_token_based_replay(log,
    petri_net, initial_marking, final_marking)
print(fitness_tbr)

precision_alignments = pm4py.precision_alignments(log,
    petri_net, initial_marking, final_marking)
print(precision_alignments)

precision_tbr = pm4py.precision_token_based_replay(log,
    petri_net, initial_marking, final_marking)
print(precision_tbr)
```

Log Filtering

Filtering an event log prior to analysis is useful for three reasons. First, it allows the analyst to focus on a subset of the log information. For example, an analyst may wish to examine all traces of the order-to-cash process for domestic customers, or for business customers. Or an analyst may wish to examine only those cases that show some compliance problem.

Second, filtering allows the analyst to split the event log in order to identify differences or similarities. For example, filtering the log of the order-to-cash process for domestic customers allows the analyst to identify differences in how domestic and overseas customer orders are processed.

Third, filtering simplifies automatically discovered models. Many automatic discovery algorithms produce very complex models when the actual processes are complex or when there is a large amount of variation or noise in the event log. Noise does not necessarily mean invalid data, but data that appears infrequently or could be considered very atypical. Such noise, when included in the event log for process discovery, can "clutter up" the resulting model, making it difficult or impossible to understand.

PM4Py provides a number of different filters, with a few examples shown in Table 8.1. Information on other filters can be found on the PM4Py website.

<code>filter_activities_rework</code>	Keep cases where the specified activity occurs at least n times
<code>filter_case_size</code>	Keep cases having a length between n and m events
<code>filter_case_performance</code>	Keep cases having a duration between n and m seconds
<code>filter_directly_follows_relation</code>	Keep cases where A is followed immediately by B
<code>filter_end_activities</code>	Keep cases that end with the specified activity
<code>filter_event_attribute_values</code>	Keep cases or events in cases that satisfy the specified condition
<code>filter_eventually_follows_relation</code>	Keep cases where A is eventually followed by B
<code>filter_start_activities</code>	Keep cases that start with the specified activity
<code>filter_time_range</code>	Keep events occurring between two timestamps
<code>filter_trace_attribute_values</code>	Keep cases that satisfy the specified condition

Table 8.1: Example event log filter functions in PM4Py

Hands-On Exercises – Basic Log Information

The following exercises are designed to be used with the event log in the running examples above. Tips are provided to guide you and to provide links to the documentation for important functions to use.

1. What are the different types of activities in the log?
 - Use the Pandas `unique()` function
2. How often does each activity occur in the log?
 - Use the Pandas `value_counts()` function
3. Filter the log for complete cases, that is, retain only those cases that end with activity "Pay invoice".
 - Use `pm4py.filtering.filter_end_activities`
4. Plot the case durations for the complete cases. What do you notice?
 - Use `pm4py.stats.get_all_case_durations`
 - Put case durations into a `pd.DataFrame`
 - 1 day = 86400 seconds
 - Use `px.histogram` or `pm4py.vis.view_case_duration_graph`
5. What is the mean case duration?
 - Use the Pandas `mean()` function on the result of the previous exercise

Hands-On Exercises for PM4Py – Automatic Process Discovery

The following exercises are designed to be used with the event log in the running examples above. Tips are provided to guide you and to provide links to the documentation for important functions to use.

1. Using the mean case duration identified in the previous exercise, split the log on the mean case duration; that is, one sub-log should contain traces that are shorter than the mean, the other sub-log should contain cases that take longer than the mean.
 - Use `pm4py.filtering.filter_case_performance`
2. Discover BPMN models for each partial log and compare them. How do they differ?
3. Discover a BPMN model from the total log. How does it differ from the models discovered for the partial logs?
4. Calculate and compare the fitness and precision values of the models discovered from the partial log and the total log.

Hands-On Exercises for PM4Py – Performance Analysis

The following exercises are designed to be used with the event log in the running examples above. Tips are provided to guide you and to provide links to the documentation for important functions to use.

1. What is the activity with the longest mean time? Activities taking a long time may be a bottleneck in the process flow.
 - Create a new column as the difference between the 'Complete Timestamp' and 'start_timestamp' columns.
 - Use the Pandas `groupby()` and `mean()` functions to group the data frame by activity.
2. What is the mean number of activities for each case? Long cases with many activities may indicate problems or overly complex processes.
 - Calculate the number of activities for each case using the Pandas `groupby()` and `count()` functions on the dataframe
3. Which activities are carried out more than once for some case? Repeated activities may indicate re-work or fixing of mistakes.
 - Calculate the number of instances for each case for each activity using the Pandas `groupby()` and `count()` functions on the dataframe

Hands-On Exercises for PM4Py – Conformance Analysis

The following exercises are designed to be used with the event log in the running examples above. Tips are provided to guide you and to provide links to the documentation for important functions to use.

1. Are there cases that contain activity "Pay invoice" but do not contain activity "Send invoice"? Non-compliant cases may represent a problem with controls and compliance.
 - Use [filter_eventually_follows_relationship](#)

8.7 Performance Mining

Performance mining is that aspect of process analytics that analyzes the temporal performance of a process. Information of interest are the durations of the activities (*service time*), the *waiting times* between activities, and the overall case durations. For each of those, different summary statistics are useful, such as the mean, median, standard deviation, maximum and minimum.

The easiest and most interpretable way to do this in PM4Py is to annotate the DFG with this information. For example, the following Python code block calculates the median waiting times between activities and shows them in the DFG. The annotated DFG is shown in Figure 8.8. Because the example event log did not contain start and end times for each activity, it is not possible to show the service times on the DFG, that is, the duration of the activities.

```
perf_dfg, start_activities, end_activities = \
    pm4py.discover_performance_dfg(log)

pm4py.view_performance_dfg(perf_dfg,
    start_activities, end_activities,
    aggregation_measure='median')

pm4py.save_vis_performance_dfg(perf_dfg,
    start_activities, end_activities,
    file_path='perfdgf.png', rankdir='TB')
```

This example shows a few problems with the process. For example, the median wait time between creating a request for quotation and analyzing it is 6 days. If the request needs to be amended, it then has to wait another 8 to 10 days to be analyzed again. These long wait times indicate performance problems in the process. This may stem from a lack of resources, or insufficient prioritization, or other process issues, that needs be investigated in detail by the process analyst.

Another useful tool in process performance analysis is the *dotted chart*. An example dotted chart is shown in Figure 8.9. The horizontal axis represents the time stamp of each activity or event, while the vertical axis represents the case ID – each row in the

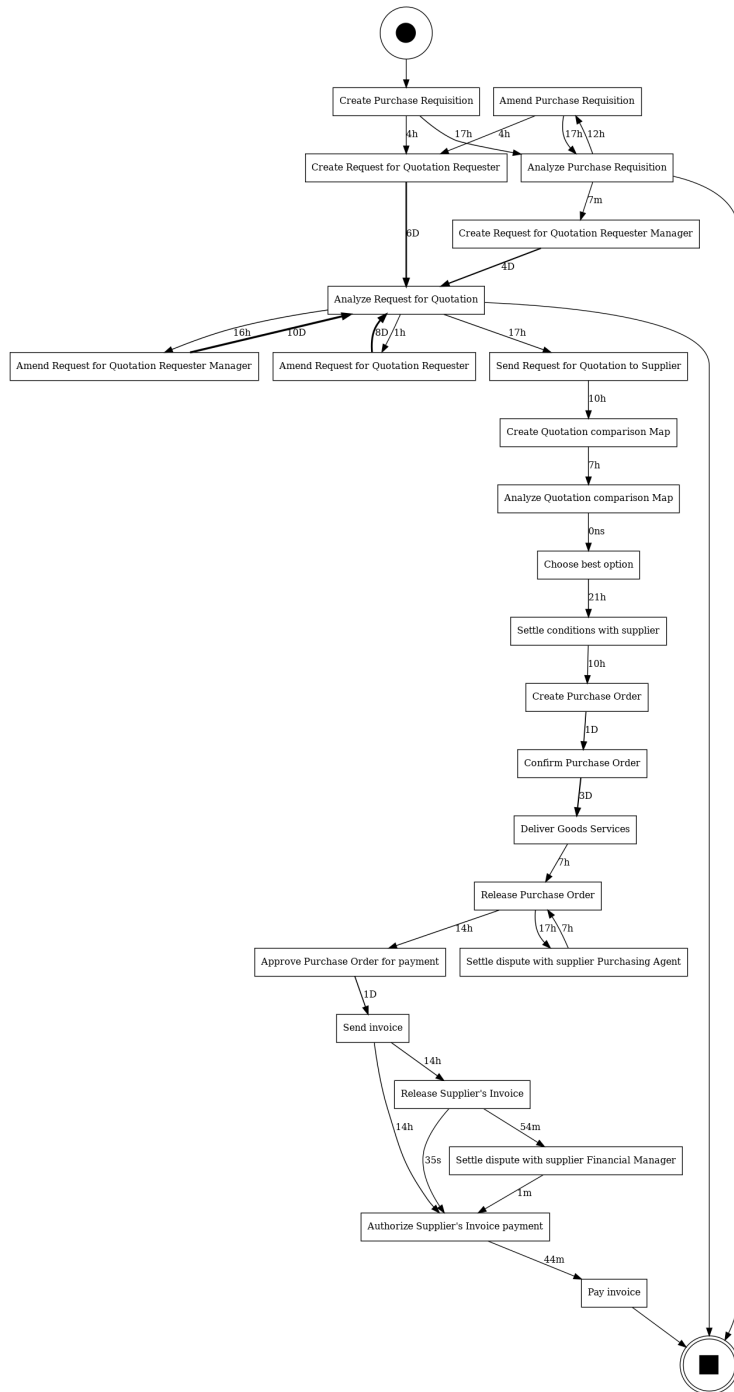


Figure 8.8: DFG annotated with median waiting times

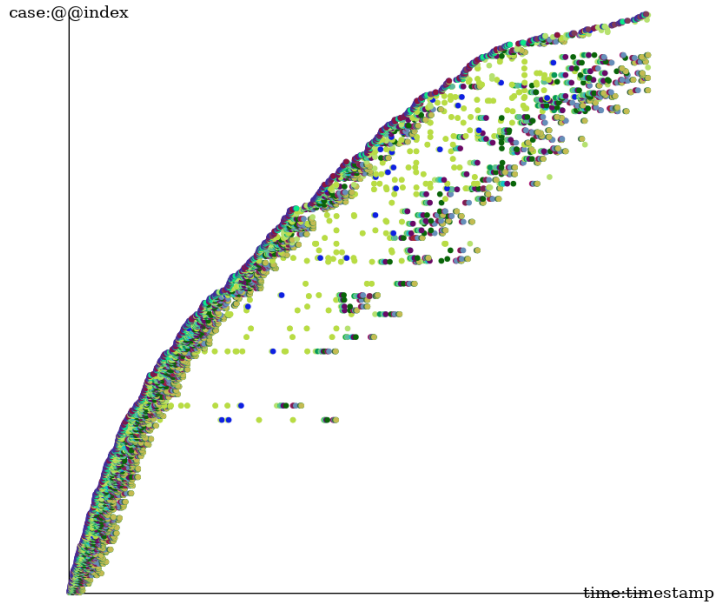


Figure 8.9: Example of a Dotted Chart

diagram contains the events of one trace. The colors correspond to different types of activities. The following Python code produces the example in Figure 8.9.

```
perf_dfg, start_activities, end_activities = \
    pm4py.discover_performance_dfg(log)

pm4py.view_performance_dfg(perf_dfg,
    start_activities, end_activities,
    aggregation_measure='median')

pm4py.save_vis_performance_dfg(perf_dfg,
    start_activities, end_activities,
    file_path='perfdg.png', rankdir='TB')
```

A dotted chart can be used to show batching of activities, that is, activities in different cases that are not spread out in time but are executed at the same time. This may indicate that some cases have to wait for the next batch to be processed, leading to potential delays. A dotted chart can also show different variants, by visually highlighting different types of performance. In the example of Figure 8.9, it is clear that many cases are finished quickly, while others take a long time. In particular, the cases that arrive early or very late in the log tend to be those that finish quickly.

A dotted chart can also be useful to examine the case arrival rates. In the example in Figure 8.9 it is clear that early in time, cases arrive more frequently than later in time

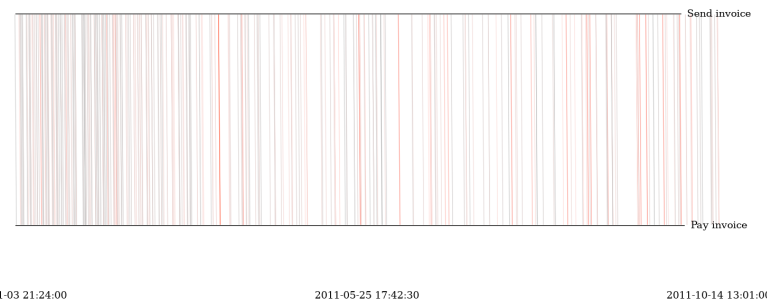


Figure 8.10: Example of a Performance Spectrum

(the curve is steeper there). This may indicate a shift in the demand for a particular product or service.

A performance spectrum graph, like the one in Figure 8.10 shows the waiting times between two activities, the one at the top and the one at the bottom. The horizontal axis represents time. Lines that are slanted or run diagonally at an angle indicate a long waiting time. A performance spectrum can also show when lines are "bunched together" or "sparsely distributed", indicating variations in the rate at which one activity finishes or the next activity begins. The following Python code block produces the graph in Figure 8.9.

```
pm4py.view_performance_spectrum(log,
    ['Send invoice', 'Pay invoice'])

pm4py.save_vis_performance_spectrum(log,
    ['Send invoice', 'Pay invoice'],
    'perfспектum.png')
```

To indicate when a process that delivers a service is in high demand and requires high capacity, the distribution of events over time should be considered. This can be done by day-of-the-week, by month-of-the-year, or by week-of-the-year, as the following PM4Py functions show. An example of event distribution by week-of-the-year is shown in Figure 8.11.

```
pm4py.view_events_distribution_graph(log, 'days_week')
pm4py.view_events_distribution_graph(log, 'days_month')
pm4py.view_events_distribution_graph(log, 'months')
pm4py.view_events_distribution_graph(log, 'weeks')
```

The example in Figure 8.11 shows that this process is very busy early in the year, in approximately the first quarter, but not at all in the last quarter of the year, except for some activity in the final week of the year. This uneven demand requires adequate

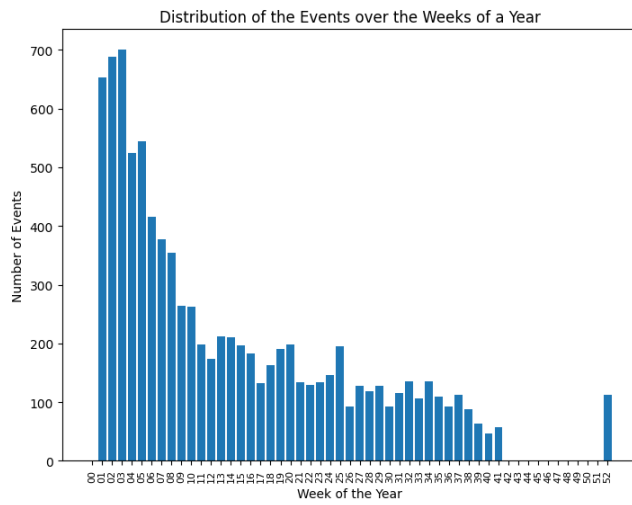


Figure 8.11: Event distribution over time

capacity planning over of the organization and an organization may decide to identify ways to smooth out the demand.

Plotting the events over time for the entire log as a probability density shows similar characteristics. Figure 8.12 plots the frequency (technically, a probability kernel density) of event occurrence, produced by the following Python code.

```
pm4py.view_events_per_time_graph(log)
pm4py.save_vis_events_per_time_graph(log, 'eventspertime.png')
```

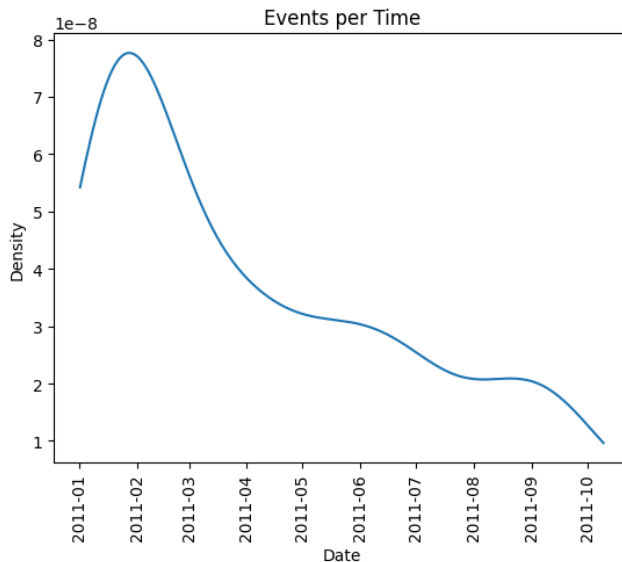


Figure 8.12: Events per time graph

8.8 Organizational Mining

Organizational mining focuses on the resources and their roles in a process. It uses the data in an event log to identify how resources and roles work together to execute process instances.

A simple way to focus on organizational roles or resources is to construct a DFG, but using the resource or role information for the nodes of the graph, instead of the activity names. The DFG then expresses how often one resource or role follows another in the execution of the cases; in other words, how often one resource or role passes work to another (or to itself). The following Python code block produces the *handover-of-work network* shown in Figure 8.13. Notice how the same PM4Py function for the DFG discovery is used, but a different data frame column is specified for the "activity_key" parameter.

```
dfg, start, end = pm4py.discover_dfg(log, activity_key='Role')

pm4py.view_dfg(dfg, start, end, rankdir='LR')

pm4py.save_vis_dfg(dfg=dfg,
    start_activities=start,
    end_activities=end,
    file_path='handover.png', rankdir='TB')
```

A similar analysis can be performed using the PM4Py function

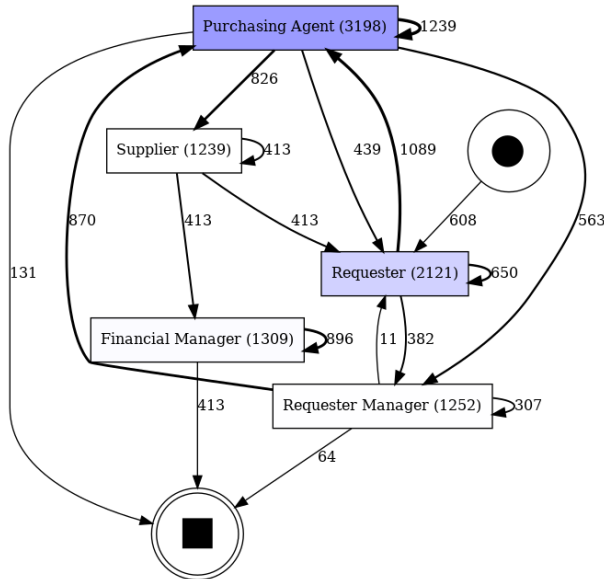


Figure 8.13: Example handover-of-work network

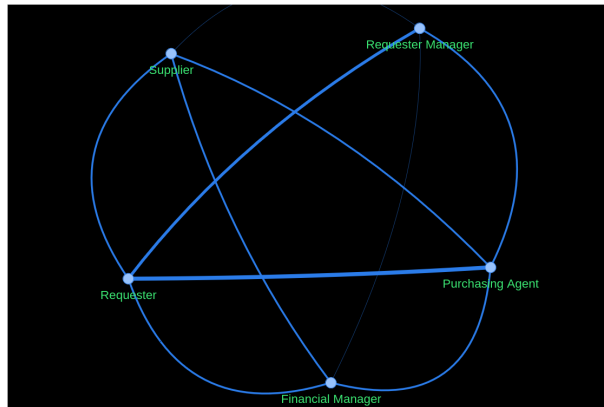


Figure 8.14: Example working-together network

`discover_working_together_network()`, as shown in the following Python code and in Figure 8.14. Note that this graph is normally interactive.

```
sna_graph = pm4py.discover_working_together_network(log,
    resource_key='Role')
pm4py.view_sna(sna_graph, variant_str='pyvis')
pm4py.view_sna(sna_graph, variant_str='networkx')
```

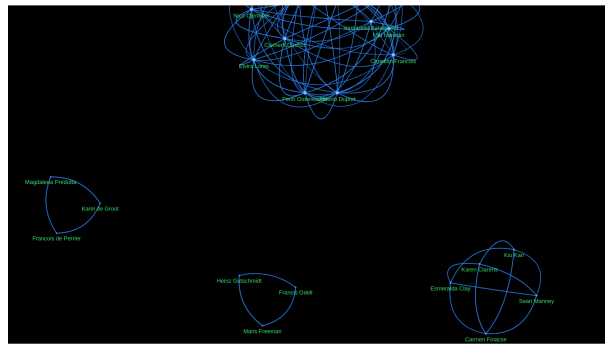


Figure 8.15: Activity-based resource similarity graph

Yet another analysis focuses on identifying resources that perform the same sets of activities. This is useful for identifying implicit roles or resources with similar skills.

```
roles = pm4py.discover_organizational_roles(log)
print(roles)
```

Another way to achieve this goal is with the following Python code, which produces the activity-based resource similarity graph in Figure 8.15. The graph shows four clusters of resources that are similar to each other within their cluster, i.e. they perform similar sets of activities in this process.

```
sna_graph = pm4py.discover_activity_based_resource_similarity(log)

pm4py.view_sna(sna_graph, variant_str='networkx')
pm4py.view_sna(sna_graph, variant_str='pyvis')

pm4py.save_vis_sna(sna_graph, 'ressimilarity.png',
variant_str='networkx')
```

8.9 Review Questions

The following review questions help you evaluate your understanding of this material.

1. What is business process analytics and why is it important in improving operational efficiencies?
2. Define a business process. What are the typical elements that constitute a business process?
3. What roles do resources play in a business process? Provide examples.
4. What is the Business Process Modeling Notation (BPMN)? Why is it widely used?

5. Explain the symbols used in BPMN to represent events, activities, gateways, and dependencies.
6. Describe the function of an exclusive gateway in a BPMN diagram. Provide an example from the given material.
7. Explain what a parallel gateway is and provide an example of how it is used in business process modeling.
8. What is an inclusive gateway? Give an example of its application in a business process.
9. Given a simple process scenario, draw a BPMN diagram using appropriate symbols for events, activities, and gateways.
10. Define a "case" in the context of a business process. Provide an example.
11. What is a "trace"? How does it relate to a case?
12. Explain the term "event" in process mining. What might an event signify in a business process event log?
13. Describe the typical lifecycle of an activity in a business process. Refer to the XES standard lifecycle model.
14. What are some possible states and transitions an activity might go through during its lifecycle?
15. Differentiate between "event attributes" and "case attributes." Provide examples of each.
16. Why is it important to associate events with resources and timestamps?
17. Define an "event log." What kind of information does it typically contain?
18. What is the XES file format?
19. How does a CSV file format differ from an XES file when used for storing event log data?
20. Explain what is meant by automated process discovery in process analytics. Why is it considered a crucial initial step for many organizations?
21. Define conformance checking. What are its typical uses in process analytics?
22. Discuss how conformance checking can demonstrate compliance with normative process models or business rules during audits.
23. Describe what is meant by performance mining in the context of process analytics.
24. Identify and explain the types of problems that performance mining can help to uncover within a process.
25. What is variants analysis, and why might it be useful for organizations that operate in multiple locations or business units?
26. Provide examples of insights that can be gained from performing variants analysis on business processes.
27. Explain the concept of process prediction and its importance in process analytics.
28. Discuss potential applications and provide examples of process prediction in managing business processes and customer relations.

The following questions are specific to PM4Py but the main concepts apply to other process analytics software tools as well.

29. Describe the role of the following columns in the process analytics context and why they need to be specified:

- Case identifier
 - Activity name
 - Timestamp
 - Resource
30. How does PM4Py determine which columns represent the case ID, activity name, timestamp, and resource in the event log data? Include a brief explanation of how these columns are transformed or defined in the provided Python code.
31. Compare and contrast the processes of loading event logs from CSV files and XES files into PM4Py. Discuss the advantages and disadvantages of using each file format.
32. What Python code would you use to calculate the number of cases and the number of events in an event log? Explain what each line of code does.
33. Describe the purpose of the following PM4Py functions:
- `get_start_activities()`
 - `get_end_activities()`
 - `get_all_case_durations()`
34. How does the `split_by_process_variant()` function in PM4Py work?
Describe what it returns and how these returns can be used in process analysis.
35. Explain what a directly-follows graph (DFG) is and its importance in process discovery.
36. Discuss the observations that can be made from a DFG (e.g., start activities, end activities, loops, and possible re-work).
37. Compare the Inductive Miner and Heuristics Net Miner provided by PM4Py in terms of how they process a DFG to discover BPMN models.
38. Define the four aspects of model quality in process mining.
39. Describe how token-based replay and alignment-based fitness methods work to assess the fitness of a process model.
40. Explain how precision differs from fitness in the context of process model quality and why both are important.
41. Explain the importance of filtering an event log before conducting process analysis. Include three reasons why filtering might be necessary.
42. Describe how filtering can help an analyst focus on specific aspects of a process. Provide examples of different subsets an analyst might focus on within an event log.
43. What types of filters might an analyst use to refine an event log before analysis? Provide examples or scenarios where specific filters would be particularly useful.
44. Explain what performance mining in process analytics entails and why it is important.
45. Discuss the types of information typically analyzed in performance mining, such as service time, waiting times, and overall case durations. Why are these metrics important?
46. Review the purpose of a dotted chart in performance analysis. How does this visualization help in identifying batching of activities or case arrival rates?
47. Describe what a performance spectrum graph shows and how it can be used to identify performance issues between two activities.
48. How can the distribution of events over time be used to inform capacity planning

in an organization?

49. Consider the tools and methods described (DFG, dotted chart, performance spectrum, event distribution graphs). Discuss how each can contribute to a comprehensive performance analysis of a business process.
50. Explain the concept of organizational mining and its importance in understanding process execution within an organization.
51. Describe what a handover-of-work or working-together network is. What does this type of network reveal about the interactions between roles or resources?
52. Review the usefulness of analyzing resources that perform similar sets of activities. How does identifying implicit roles or skills similarity benefit an organization?

Chapter 9

Introduction to Supervised Machine Learning

Sources and Further Reading

The material in this chapter is based on the following sources. They are freely available. Consult them for additional information.

Gareth James, Daniel Witten, Trevor Hastie and Robert Tibshirani: *An Introduction to Statistical Learning with Applications in R*. 2nd edition, corrected printing, June 2023. (ISLR2)

<https://www.statlearning.com>

Chapters 2, 3, 4, 5

The book by James et al. provides an easy introduction to machine learning at the introductory undergraduate level. It focuses on applications, not mathematics, and contains many exercises using R. Concepts are well explained and illustrated. There is a similar book available by the same authors with applications in Python.

Trevor Hastie, Robert Tibshirani, and Jerome Friedman: *The Elements of Statistical Learning*. 2nd edition, 12th corrected printing, 2017. (ESL)

<https://hastie.su.domains/ElemStatLearn/>

Chapters 2, 3, 4, 7

The book by Hastie et al. still sets the standard for statistical learning. It is widely used and cited. Its treatment is more technical than the previous book and there are no exercises in R or Python. However, it covers the concepts in more depth (and a few more formulas). However, it is still very accessible even to an undergraduate audience.

Kevin P. Murphy: *Probabilistic Machine Learning – An Introduction*. MIT Press 2022.

<https://probml.github.io/pml-book/book1.html>

Chapters 4, 6, 9, 10, 11

Murphy's book is available under a creative commons license. It is a somewhat more technical treatment of the material, but with many illustrations and examples. It is quite comprehensive in its coverage and targeted at the advanced undergraduate or graduate student.

9.1 Introduction

Supervised machine learning is the training or fitting of statistical models for prediction tasks when the correct target outcome is known. It is called "training" because we train a statistical model to make predictions for future observations based on past data. It is also called "fitting" because, for parametric models, that is models with parameters, we adjust the model parameters to ensure the model output is a good fit with the known, correct target outcome; that is, we fit the model to the data.

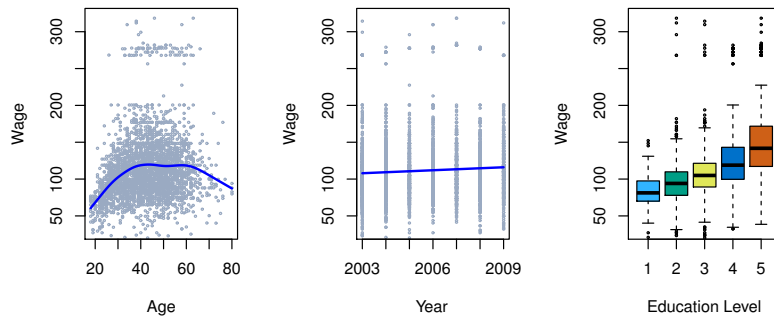
Depending on the application area and research discipline, different terminology may be used. In this chapter, the term *inputs* is used for variables used to make a prediction, and the term *outputs* is used for the predicted variables. Other terms for inputs are *predictors*, *independent variables*, and *features* although there are fine but important distinctions that are covered later. Other terms for outputs are *targets*, *responses*, and *dependent variables*.

Many methods in supervised learning are *parametric methods* that assume a functional relationship of the form

$$y = f(x) + \epsilon$$

where the function f is approximated by a function \hat{f} that is characterized by a set of parameters. The values for these parameters are learned or estimated in order to optimally fit the model to the existing training data. Once the optimal parameters are estimated, the fitted or trained model can be used to predict the output for new inputs.

On the other hand, *non-parametric methods* do not assume a functional form. They can therefore be more flexible.



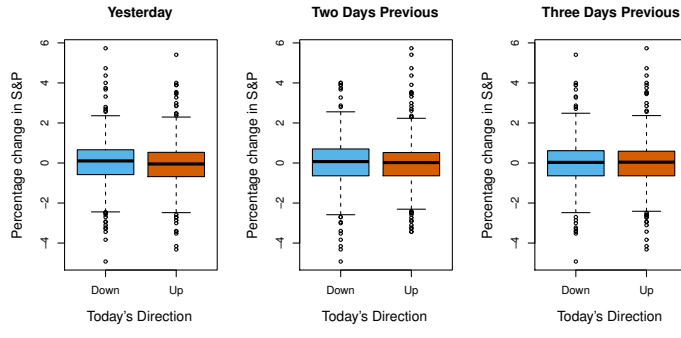
Source: ISLR2 Figure 1.1

Figure 9.1: Example regression model

Depending on the type of output, one distinguishes regression from classification. In *regression analysis*, the output is quantitative, typically real-valued. The quality of a model is measured by the numeric differences between actual and predicted output. Figure 9.1 shows an example of a regression model that predicts the output "Wage" from the inputs "Age", "Year", and "Education Level". Note that one of the inputs is quantitative while the other two are categorical (although both "Year" and "Education Level" could conceivably be treated also as quantitative in another model).

There are many different regression metrics, some parametric, some non-parametric. The highlighted entries in the list are covered in this and the following chapter; later chapters will also cover neural network regression and regression trees.

- **Parametric Methods**
 - **Linear Regression**
 - **Ridge and Lasso Regression**
 - Principal components regression
 - Non-linear regression
 - Neural network regression
- **Non-Parametric Methods**
 - **K-Nearest-Neighbours (KNN)**
 - Regression trees
 - Smoothing splines
 - Multivariate adaptive regression splines
 - Kernel regression



Source: ISLR2 Figure 1.2

Figure 9.2: Example classification model

In contrast, the output of *classification* is categorical or qualitative. The quality of a model is measured by the proportion of correct classifications (accuracy) and related metrics. Figure 9.2 shows an example of a classification model where the binary output "Today's Direction" (of the stock market) is to be predicted from the inputs "Yesterday", "Two Days Previous", and "Three Days Previous". Note that all three inputs are qualitative, although quantitative inputs can also be used in classification.

There are many different classification methods, most of which are parametric, except for decision trees and KNN. The highlighted methods are covered in this and the following chapter; decision trees and neural networks are covered in later chapters.

- Decision trees
- Random forests
- Bayesian networks
- Support vector machines
- Neural networks
- **Logistic regression**
- **Naive Bayes**
- Probit model
- Genetic programming
- **K-Nearest-Neighbours (KNN)**

9.2 Explanation and Prediction

Explanation and *prediction* both use statistical models. However, they differ in their goals and methods. *Explanation* seeks to understand the *causal mechanisms* in the

Explanation	Prediction
Causation	Association
Theory	Data
Retrospective	Prospective
Bias	Variance

Based on: Shmueli, G. (2010). To Explain or To Predict?. *Statistical Science*, 25, 289-310.

Table 9.1: Differences between explanation and prediction

world, that is, understand why a particular output is observed for a given input. The statistical model is intended to represent, or be isomorphic to (have the same form as), the causal processes. Explanation is often concerned with *theory testing*. Model parameters are assumed to represent the strength of a causal effect hypothesized by some theory. Scientists use a *representative sample* of observations to *infer* the value of a parameter in the larger population in order to support or reject a causal theory. Explanation aims for relatively simple models, e.g. linear ones, that can be understood and interpreted by humans. Explanation is *retrospective*, that is, backward looking. It uses the observed data in the sample to fit a model, but does not normally collect additional data to further verify the fit of that model on other data. In other words, explanation is concerned with most closely fitting a model to a single sample, which is assumed to be representative of the population. This is called "bias minimization", a term explained in more depth later.

In contrast, prediction is not concerned with causal processes or with models that represent or are isomorphic to causal relationships in the world. A predictive model that produces accurate predictions is satisfactory even if it does not represent the true causal relationship. In other words, prediction is concerned with *association*, not with causation. In contrast to explanation, prediction does not know the concepts of population, sample, and inference from sample values to population values. Instead, prediction uses the term "training data" and requires that the training data be representative of future observations for which predictions are to be made, not of some larger population. In that sense, prediction is *prospective*, that is, forward looking. Prediction focuses on individual observations, rather than the fit of the entire model. Moreover, because the models are not intended to represent causal relationships and theories, they may be more complex and need not be humanly understandable or interpretable. Instead of focusing on fitting a statistical model to a single set of observations, as is done in explanation, prediction recognizes the variability that is introduced by different sets of training data. This is termed variance, a concept explained in more depth later. Prediction tries to minimize both the bias and the variance in order to learn models that accurately predict future, yet unseen observations.

Table 9.1 provides a summary of the differences between explanation and prediction.

Hands-On Exercise

For each of the following problems, decide if it is a prediction or inference/explanation problem:

1. How do real estate prices vary with location and age?
2. What is the most important predictor of real estate prices?
3. What is the expected sales price for a house at 310 Elizabeth Ave?
4. Is the month of the sale an important predictor of real estate prices?
5. Calculate the difference in expected sales prices for the house at 310 Elizabeth Avenue when sold in August and February
6. When should a house be sold to achieve the best price?

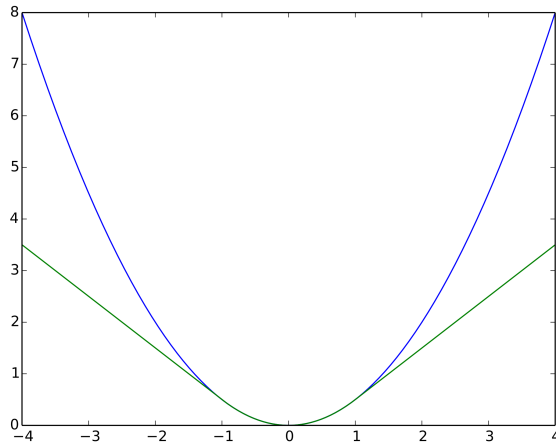
9.3 Bias and Variance in Regression Analysis

The predictive quality of a regression model is typically evaluated by the *mean squared error* (MSE) or the *mean absolute error* (MAE). The error of the model in predicting the correct output, that is, the difference between prediction and true output, is often called the *loss function*, which is to be minimized for optimal fit. The mean absolute error is sometimes preferred because it is more robust to the presence of outliers as it does not square the difference between prediction and target; other *loss functions* are the *mean absolute percentage error* or the *Huber loss*. The Huber loss function, shown in Figure 9.3, combines a square error for small values with an absolute error for larger values, making it also robust to outliers.

$$\begin{aligned} \text{MSE} &= \frac{1}{n} \sum_{i=1}^n (y_i - \hat{f}(x_i))^2 \\ \text{MAE} &= \frac{1}{n} \sum_{i=1}^n |y_i - \hat{f}(x_i)| \\ \text{MAPE} &= \frac{1}{n} \sum_{i=1}^n \left| \frac{y_i - \hat{f}(x_i)}{y_i} \right| \\ L_{\text{Huber}} &= \begin{cases} \frac{1}{2}(y - \hat{f}(x))^2 & \text{for } |y - \hat{f}(x)| \leq \delta \\ \delta(|y - \hat{f}(x)| - \frac{1}{2}\delta) & \text{otherwise} \end{cases} \end{aligned}$$

The parameter δ for the Huber loss function can be freely chosen.

Importantly, the focus of evaluating the quality of a model should not be on the training data itself, but on how well it performs on data that were not used for training. For example, a model is trained on past stock market information, but is used to predict future stock performance; a model is trained on previous patient information, but is used to predict future patient outcomes; a model is trained on past real estate prices but is used to predict future closing prices.



https://commons.wikimedia.org/wiki/File:Huber_loss.svg

Figure 9.3: Huber loss function versus squared error

A typical strategy is then to separate *test data* from the *training data* in the form of a *holdout sample*. The model is fitted to the training data and then evaluated on the test data.

Low error on training data does NOT imply low errors on test data.

Consider the regression model shown in Figure 9.4. The left panel shows a set of observed x and y values generated from the true relationship (black line) by adding some random error. The left panel also shows three different functions that are fitted to this data. The diagonal orange line represents a simple linear regression model with only x and an intercept as predictors, that is it only has two parameters. The blue and green lines represent smoothing regression splines with different levels of flexibility.

It is evident that the orange line fits neither the observed data very well, nor is it close to the true function, the black line. It lacks sufficient *flexibility* to both approximate the data and the true model. It has been *underfitted*.

It is evident from the left panel in Figure 9.4 that the green line fits the observed data better than the blue line, but it is also clear that the green line does not fit the true model, represented by the black line, as well as the blue line does. If one were to generate another sample of observations from the black line by adding random errors, the green line is unlikely to fit this new sample very well. In other words, the green line model has been fitted to the particular characteristics, or idiosyncrasies of this one set of training data: the model has been *overfitted*.

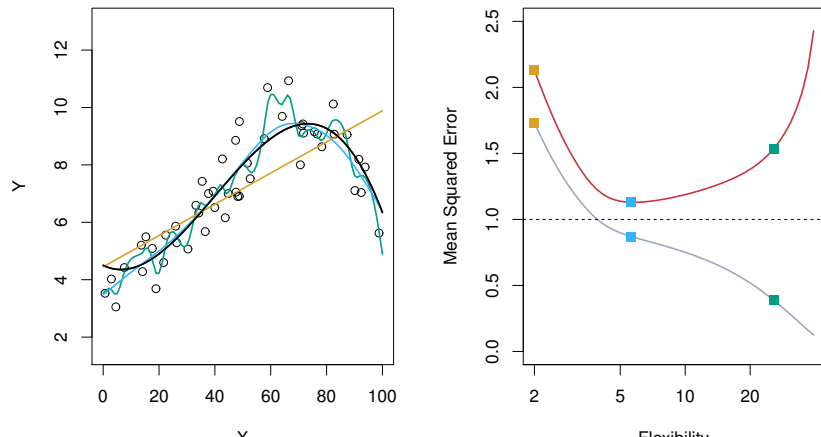
The right panel of Figure 9.4 shows the training data error (gray line) and the test data

error (red line) for the three models indicated by the coloured squares (and others in between). The underfitting orange model shows both a large training error as well as a large test error on a holdout sample. In contrast, the green overfitted model shows a very small training error but a large test error on the independent test or holdout data set. The blue model does not show the smallest training error but it does show the smallest testing error.

While Figure 9.4 has been created with non-parametric smoothing spline models, in parametric models, such as linear regression and others, the flexibility of the model to adapt itself to the training data is a function of the number of parameters that can be freely adjusted. For example, a regression model with only an intercept and the variable x as predictor will have two parameters: the slope and the intercept and is therefore less flexible than a model with intercept and predictors x , x^2 and x^3 . Figure 9.5 shows an example with parametric models. The top-left panel shows a model with predictors x and x^2 fitted to a set of training data. The bottom left panel shows a model with polynomials in x up to degree of 20. As there are only 20 data points, the model fits the data perfectly, but is unlikely to perform well on new, unseen observations. The bottom right panel in Figure 9.5 shows the train and test errors for models with different degrees of polynomials. Figure 9.5 is similar to Figure 9.4 in the essential characteristics of overfitted models, that is a low training error and a high test error.

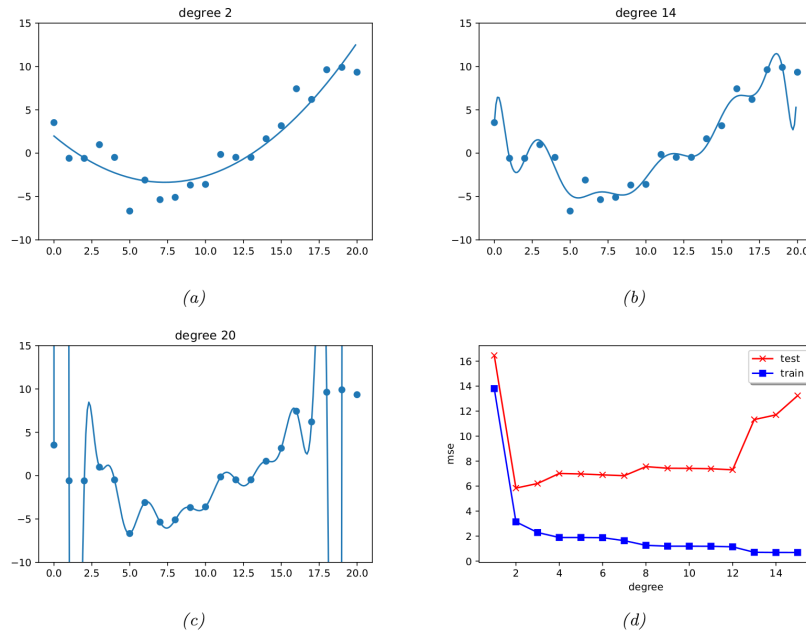
Closely related to flexibility is concept of *degrees of freedom*. In parametric models, the degrees of freedom are defined as the difference between the number of observations and the number of parameters of the model:

$$\text{DF} = n - p$$



Source: ISLR2 Fig 2.9

Figure 9.4: Fit versus flexibility of a model



Source: Murphy Figure 1.7

Figure 9.5: Fit versus number of parameters of a model

Each parameter to be estimated requires one observation and so "uses up" one degree of freedom. The model in the bottom left panel of Figure 9.5 has no degrees of freedom left, it has as many parameters p as observations n .

To better understand the concepts of underfitting and overfitting, it is useful to consider the MSE regression loss in more detail. This requires the concepts of expected value and variance of a random variable X from basic statistics.

Expected Value

$$E[X] = \sum_{i=1}^{\infty} x_i p_i \quad \text{discrete random variable}$$

$$E[X] = \int_{-\infty}^{\infty} xp(x) dx \quad \text{continuous random variable}$$

For uniform distributions or unweighted observations $p_i = p_j \forall i, j$ so that $E[X] = \frac{1}{n} \sum_{i=1}^{\infty} x_i$, i.e. expectation = mean

Variance

$$\text{Var}[X] = E[(X - E[X])^2] = E[X^2] - E[X]^2$$

For zero-centered variables $E[X] = 0$ so that $\text{Var}[X] = E[X^2]$

Equipped with these concepts, one can rewrite the MSE to decompose it into three parts:

$$\begin{aligned}\text{MSE} &= E[(y - \hat{f})^2] \\ &= E[y^2 - 2y\hat{f} + \hat{f}^2] \\ &= E[y^2] - 2E[y\hat{f}] + E[\hat{f}^2]\end{aligned}$$

Using the definitions for expected value and variance, each of part of the MSE can be further rewritten:

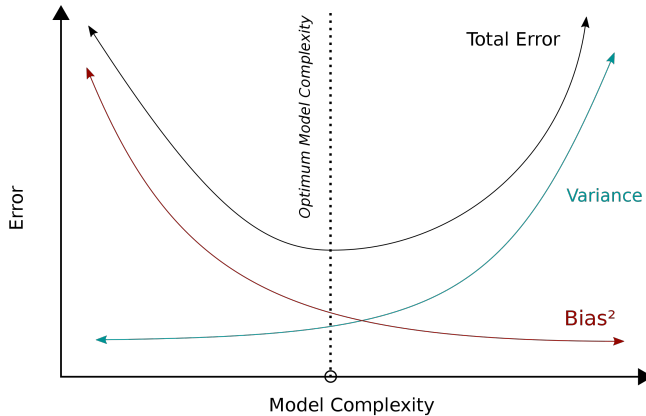
$$\begin{aligned}E[\hat{f}^2] &= E[\hat{f}^2] - E[\hat{f}]^2 + E[\hat{f}]^2 \\ &= \text{Var}[\hat{f}] + E[\hat{f}]^2\end{aligned}$$

$$\begin{aligned}E[y^2] &= E[(f + \epsilon)^2] \\ &= E[f^2] + 2E[f\epsilon] + E[\epsilon^2] \\ &= f^2 + 2f \cdot 0 + \sigma^2 \quad (f \text{ is not random and } E[\epsilon] = 0) \\ &= f^2 + \sigma^2\end{aligned}$$

$$\begin{aligned}E[y\hat{f}] &= E[(f + \epsilon)\hat{f}] \\ &= E[f\hat{f}] + E[\epsilon\hat{f}] \\ &= E[f\hat{f}] + E[\epsilon]E[\hat{f}] \\ &= E[f\hat{f}] + 0 \cdot E[\hat{f}] \\ &= fE[\hat{f}]\end{aligned}$$

Putting this all together, we can rewrite the MSE as follows:

$$\begin{aligned}\text{MSE} &= f^2 + \sigma^2 - 2fE[\hat{f}] + \text{Var}[\hat{f}] + E[\hat{f}]^2 \\ &= (f - E[\hat{f}])^2 + \sigma^2 + \text{Var}[\hat{f}] \\ &= \text{Bias}[\hat{f}]^2 + \sigma^2 + \text{Var}[\hat{f}]\end{aligned}$$



[https://commons.wikimedia.org/wiki/File: Bias_and_variance_contributing_to_total_error.svg](https://commons.wikimedia.org/wiki/File:Bias_and_variance_contributing_to_total_error.svg)

Figure 9.6: Bias and variance trade-off

The *bias* of an estimated model \hat{f} expresses how far the expected value of the estimated model \hat{f} is away from the true target value f , while the *variance* of an estimated model \hat{f} expresses how much the expected value of the estimated model varies with different inputs or data sets. The final part of the MSE is the *irreducible error* σ^2 that represents the random variations of the data around the true target value f .

An underfitting model will necessarily have a large bias as it is not close to the true model. This was illustrated by the orange line model in Figure 9.4. As underfitting models are often models that are too simple, they tend to have small variance, but this is not necessarily true for all models.

In contrast, overfitting models necessarily have a large variance. Because they are fitted to the idiosyncratic specific values of the training data, they do not generalize well to new, unseen test data. New data, even if it is drawn from the same probability distribution or population, will lead to very different predicted outputs. Thus, a large variance of a model is manifested by a large test error. A severely overfitted model may also have a large bias. An example of this is the overfitted green model in Figure 9.4, or the degree 20 polynomial model in Figure 9.5. Neither of these are close to the true model.

A well-fitting model is one that does not minimize the bias or the variance but finds the overall optimum by minimizing the joint error. To estimate the variance, it is necessary to apply the model to independent test data. In other words, a well-fitting model is one that minimizes the test error. This central idea is known as the *bias and variance trade-off* and is illustrated in Figure 9.6. Note that the total error also includes the *irreducible error* which cannot be removed but is independent of model complexity.

Returning to the characteristics of explanation and prediction, it is now clear that explanation aims to minimize only the bias by seeking to identify the true model. In

contrast, prediction also includes a reduction of variance and focuses on minimizing the overall or joint error.

While this section has derived the concepts of bias and variance using regression models, those concepts also apply to classification models. However, the classification loss functions do not lend themselves to the easy separation of the error as the MSE loss function above.

9.4 Model Quality in Classification

In classification models, the primary quality criterion is the *error rate*, which can be calculated both for training and for test data as the proportion where the predicted class \hat{y}_i is not the true class y_i :

$$\frac{1}{n} \sum_{i=1}^n I(y_i \neq \hat{y}_i)$$

Here, $I(\cdot)$ is the *identity function* that is 1 if its argument is true, 0 otherwise.

Classification methods typically produce as output the probability of an observation belonging to any of k classes. A decision rule is then required to actually classify an observation based on this probability. A *Bayes classifier* assigns each observation to the class j with the highest probability, given its predictor values x_0 :

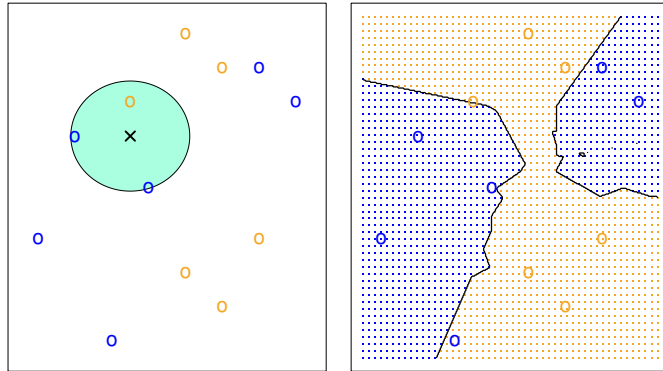
$$\operatorname{argmax}_j \Pr(Y = j | X = x_0)$$

Consequently, the error rate can be written as:

$$1 - E \left(\operatorname{argmax}_j \Pr(Y = j | X) \right)$$

The most common type of classification is *binary classification*, which assigns observations to one of two classes, e.g. true or false, normal or abnormal, good or bad, zero or one, etc. *Multinomial classification*, also called *multi-class classification* assigns observations to one of more than two classes. This section first considers binary classification before moving to multinomial classification.

A Bayes classifier is an *ideal classifier*. The Bayes classifier error rate is the irreducible error and forms the lower bound of practically achievable classification error rates. The Bayes classifier is an ideal classifier because in practice the probabilities of class membership, conditional on the observed predictor values, are unknown and must be estimated from data using a statistical model. However, estimation introduces error.



Source: ISLR2 Figure 2.14

Figure 9.7: KNN example for binary classification

A simple, non-parametric classifier is the k-Nearest Neighbour (KNN) method. This classifier identifies a set of k observations closest to an observation x_0 whose class is to be predicted, called the neighbourhood N_0 . The class membership probabilities are then estimated as the proportions of observations in the neighbourhood that belong to the different classes j :

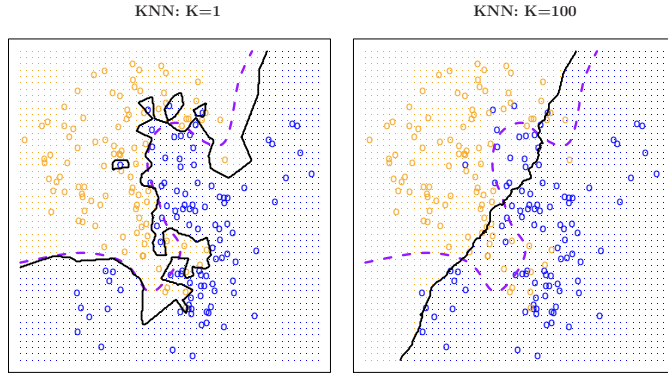
$$\Pr(Y = j | X = x_0) = \frac{1}{K} \sum_{i \in N_0} I(y_i = j)$$

Here, $I(\cdot)$ is the identity function that is 1 if its argument is true, and 0 otherwise.

Figure 9.7 provides an example for $k = 3$. The left panel in this figure shows points for which the classes, blue or orange, are known. When predicting the class for a new point, marked by the "x", the three nearest neighbours are identified. Of these, two are blue and one is yellow. Thus, the probability of the new point being yellow is estimated as $1/3$ and that of it being blue is estimated as $2/3$. The Bayes decision rule would classify the new point as blue. The right panel shows the result of classifying a large set of points as blue or yellow. The panel clearly shows the *decision boundary* of the classifier that separates the predicted class memberships, represented by the black line.

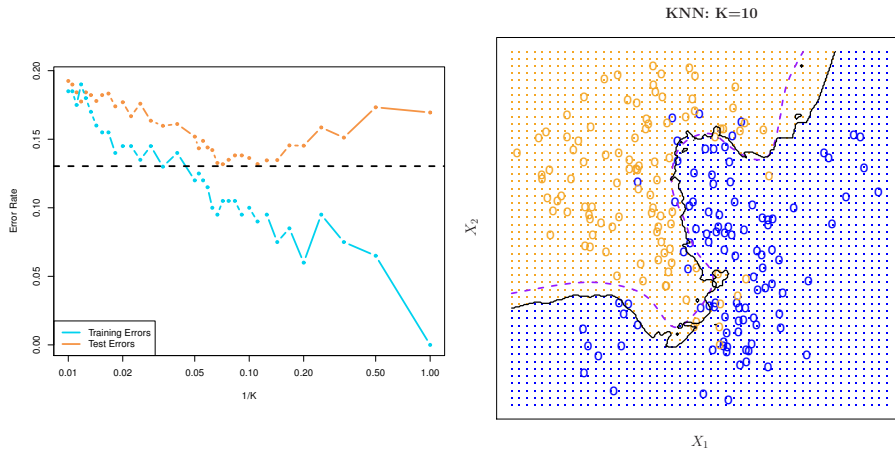
When KNN is used for regression, the predicted output value is usually estimated as the mean output values of the k neighbours in the neighbourhood N_0 .

To show how the KNN classifier behaves as a function of different values for k , consider the two panels in Figure 9.8. The left panel shows the classifications and the decision boundaries for $k = 1$. It also shows the true Bayes boundary as a dashed blue line. The KNN classifier shows relatively low bias in that its decision boundary is



Source: ISLR2 Figure 2.16

Figure 9.8: Decision boundaries of two KNN classifiers



Source: ISLR2 Figures 2.14, 2.15

Figure 9.9: KNN error rates and optimal KNN decision boundary

somewhat close to the Bayes decision boundary. However, it also shows signs of overfitting when the classifier decision boundary adapts to various individual points along both sides of the true boundary. The right panel shows a KNN classifier for the same data set with $k = 100$. It is clear that the classifier decision boundary does not follow the true decision boundary, that is, the classifier shows high bias. This classifier is underfitted. On the other hand, given that $k = 100$, it is not susceptible to individual data points or sampling changes for new data (as long as the new data is generated from the same true model), that is, it shows low variance.

In KNN classification (and also in KNN regression), the model with the lower value of k is the more flexible model, and is more likely to lead to overfitting. As k increases,

the model becomes less flexible, less likely to overfit, and increasingly more likely to underfit and have a high bias. The left panel in Figure 9.9 shows the training and test error rates for the KNN classifier for different values of k . Note the horizontal axis shows the inverse of k , that is $1/k$, larger k are to the left, smaller k are to the right in this panel. The right panel in Figure 9.9 shows the classifications and decision boundary for $k = 10$, which is close to optimal.

Hands-On Exercise The table below provides a training data set containing six observations, three predictors, and one qualitative response variable.

Obs.	X_1	X_2	X_3	Y
1	0	3	0	Blue
2	2	0	0	Blue
3	0	1	3	Blue
4	0	1	2	Yellow
5	-1	0	1	Yellow
6	-1	1	1	Blue

Suppose we wish to use this data set to make a prediction for Y when $X_1 = X_2 = X_3 = 0$ using K-nearest neighbours.

1. Compute the Euclidean distance ("l2-norm" or "Euclidean norm" of the vector difference) between each observation and the test point. The l2-norm is the root of the sum of squared differences.
2. What are your predictions for $K = 1$ and for $K = 3$? Why?
3. If the Bayes decision boundary is highly non-linear, would you expect the best value for K to be large or small? Why?

Adapted from ISLR Exercise 2.7

In binary classification, the *confusion matrix* is a useful tool for summarizing the performance of a classifier. It tabulates the number of positive and negative predictions that match the true values. The following confusion matrix shows the results of a hypothetical binary classifier for credit card defaults that assigns classes by highest probability:

$$\Pr(\text{default}=\text{Yes}|X = x) > 0.5 \text{ (Bayes)}$$

		<i>True default status</i>		Total
		No	Yes	
<i>Predicted default status</i>	No	9644	252	9896
	Yes	23	81	104
Total		9667	333	10000

Source: ISLR2 Table 4.4

The overall error rate is $\frac{23+252}{10000} = 0.0275 = 2.75\%$. However, of the true defaulters, only $81/333 = 24.3\%$ were correctly predicted. This is called the *recall* or *sensitivity*.

It also means that the error rate for the class of true defaulters is a rather high 75.7%. The overall error rate remains small mainly because there are very few true defaulters, only 333/10000. Of the non-defaulters, 9644/9667 = 99.8% are correctly predicted. This is known as the *specificity*, for an error rate of only 0.02%.

While a Bayes classification rule of choosing the highest probability class is easy to justify on probabilistic grounds, practical applications often choose a different *threshold*. In this fictitious example, credit card defaulters may pose a significant risk so that it may be worthwhile reducing the error rate for the true defaulters, even at the cost of increasing the error rate for the non-defaulters. This is shown in the following confusion matrix:

$$\Pr(\text{default}=\text{Yes}|X=x) > 0.2$$

		<i>True default status</i>		Total
		No	Yes	
<i>Predicted default status</i>	No	9432	138	9570
	Yes	235	195	430
Total		9667	333	10000

Source: ISLR2 Table 4.5

More of the true defaulters are correctly predicted and the sensitivity has improved to 58.6% while the error rate for true defaulters has been reduced to 51.4%. On the other hand, the specificity has decreased to 97.6% and the overall error rate has increased to 3.73%.

In general, a confusion matrix shows four different values, the *true negatives* (TN), *true positives* (TP), *false negatives* (FN) and *false positives* (FP) as shown in the following table:

		<i>True class</i>		Total
		No (-)	Yes (+)	
<i>Predicted class</i>	No (-)	True Neg. (TN)	False Neg. (FN)	N^*
	Yes (+)	False Pos. (FP)	True Pos. (TP)	P^*
Total		N		

From these four values, a number of model quality statistics can be computed. Frequently used are the recall, specificity, precision, accuracy, and F1 score, which are highlighted in the following list:

- Sensitivity, **Recall**, Hit Rate, True Positive Rate:

$$TPR = \frac{TP}{P} = \frac{TP}{TP + FN} = 1 - FNR$$

- **Specificity**, Selectivity, True Negative Rate:

$$TNR = \frac{TN}{N} = \frac{TN}{TN + FP} = 1 - FPR$$

- **Precision**, Positive Predictive Value:

$$PPV = \frac{TP}{TP + FP} = 1 - FDR$$

- Negative Predictive Value:

$$NPV = \frac{TN}{TN + FN} = 1 - FOR$$

- Miss Rate, False Negative Rate:

$$FNR = \frac{FN}{P} = \frac{FN}{FN + TP} = 1 - TPR$$

- Fall-out, False Positive Rate:

$$FPR = \frac{FP}{N} = \frac{FP}{FP + TN} = 1 - TNR$$

- False Discovery Rate:

$$FDR = \frac{FP}{FP + TP} = 1 - PPV$$

- False Omission Rate:

$$FOR = \frac{FN}{FN + TN} = 1 - NPV$$

- **Accuracy** (= 1 - Error Rate):

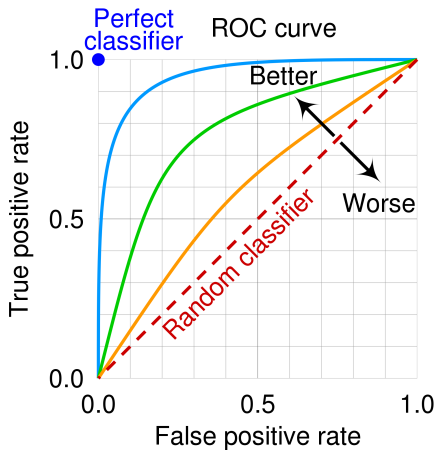
$$ACC = \frac{TP + TN}{P + N} = \frac{TP + TN}{TP + TN + FP + FN}$$

- **F1 Score** (harmonic mean of precision and recall):

$$F1 = 2 \times \frac{PPV \times TPR}{PPV + TPR} = \frac{2TP}{2TP + FP + FN}$$

- False Discovery Rate:

$$FDR = \frac{FP}{FP + TP} = 1 - PPV$$



https://commons.wikimedia.org/wiki/File:Roc_curve.svg

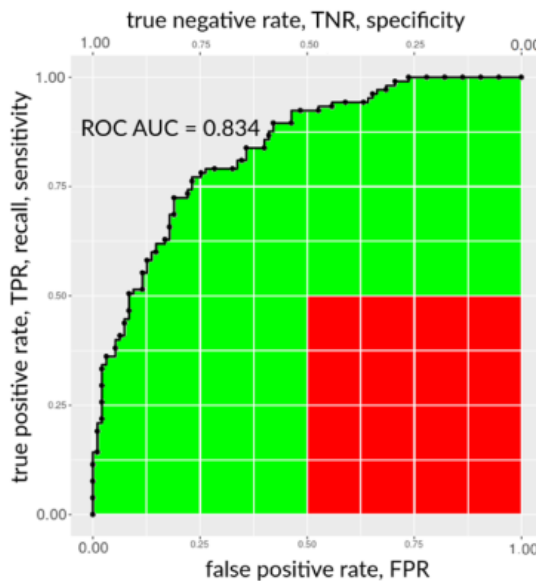
Figure 9.10: ROC curves of three example classifiers

- False Omission Rate:

$$FOR = \frac{FN}{FN + TN} = 1 - NPV$$

The above example demonstrated that the true positive rate and the false positive rate (or the true negative rate and false negative rate) are not independent of each other. Generally, increasing the true positive rate will also increase the false positive rate, because one is overall more likely to conclude that an observation is the true class (by adjusting the probability threshold). This suggests that a graph of the true positive rate against the false positive rate as shown in Figure 9.10 can summarize the classifier performance and allows the user to pick a desirable combination of true and false positive rates. The graph in Figure 9.10 is called a *Receiver Operating Characteristics* chart, or ROC chart for short. The terminology stems from early experiments with aircraft detection through radar during the 2nd world war. As shown in the figure, a random classifier is characterized by a diagonal line, and good classifiers have curves towards the upper left. Thus, the classifier shown in the green line dominates the one in the orange line, and is in turn dominated by the one represented by the blue line. All three perform better than a random classifier.

Because the ROC curve of a perfect classifier runs though the top-left corner of the ROC chart, it is natural to define the overall performance of classifier for various combinations of true and false positive rate by the area under the curve. This also allows easy comparisons of classifiers that do not dominate one another, i.e. whose ROC lines cross each other. The area-under-the-curve, or AUC for short, summarizes the classifier performance in a single number. For the classifier in Figure 9.11, the AUC is 0.834, indicated by the green and read areas. A random classifier has an AUC of 0.5 and a perfect classifier has an AUC of 1.



https://commons.wikimedia.org/wiki/File:ROC_curve_example_highlighting_sub-area_with_low_sensitivity_and_low_specificity.png

Figure 9.11: AUC of an example classifier

Hands-On Exercises

1. Consider the two confusion matrices above.
 - Compute precision and recall for the two confusion matrixes above
 - Computer accuracy and F1 values for the two confusion matrixes above
 - The two confusion matrixes above characterize two points on the ROC curve. Plot the two points for this classifier in an ROC space/diagram. Are they above or below the diagonal?
2. Consider a medical testing scenario where 1000 individuals are tested for a disease. The results are:
 - 100 people actually have the disease, and 900 do not.
 - Out of the 100 people with the disease, 90 are correctly identified as having it, but 10 are not detected.
 - Of the 900 people without the disease, 810 are correctly identified as not having it, but 90 are incorrectly identified as having the disease.

Calculate the precision, recall, sensitivity, and accuracy of the test.

Tip: Write down the confusion matrix first.

Hands-On Exercises [cont'd]

3. Given the following results from a machine learning model:
- Precision: 0.75
 - Recall: 0.60
 - Accuracy: 0.80

Answer the following questions:

- (a) What percentage of identified positives are actually positive?
- (b) What percentage of actual positives are identified by the model?
- (c) What percentage of the total classifications were correct?

4. Consider a binary classification task with the following confusion matrix at a certain threshold:
- TP: 150, FP: 50
 - FN: 30, TN: 200

Discuss how adjusting the classification threshold might affect precision, recall, and accuracy. What happens if the threshold is increased or decreased?

9.5 Multinomial Classification

Multinomial or multi-class classification assigns each observation to one of more than two classes. In this setting, the confusion matrix becomes larger, as shown in the following example, where the overall accuracy is calculated as the sum of the diagonal element divided by the sum of all elements, i.e. $\text{sum}(\text{diag}(.)) / \text{sum}(.) = 17/24 = .71$.

		True class			Prob
		0	1	2	
Predicted Class	0	4	2	0	$q_0 = 6/24 = .25$
	1	1	5	2	$q_1 = 8/24 = .33$
	2	2	0	8	$q_2 = 10/24 = .42$
Prob		p_0 $= 7/24$ $= .29$	p_1 $= 7/24$ $= .29$	p_2 $= 10/24$ $= .42$	

While the concept of a confusion matrix remains applicable, it is not immediately obvious what the true negative, true positive, false negative, and false positive values are. One way to overcome this is to reduce the classification results to a binomial case by treating each class in turn as the "positive class" and all others as "negative". This is called "one-versus-rest" (OvR), "one-versus-all" (OvA), or "one-against-all" (OaA).

In *micro-averaging*, the TP, TN, FP, and FN values are counted for all classes using OvR, and then summed. These total TP, TN, FP, and FN values are then used to compute precision, recall and other metrics. This gives equal weight to each instance but

may overemphasize the classification for a dominant majority class. Importantly, for micro-averaging, precision equals recall equals accuracy.

In *macro-averaging*, the TP, TN, FP, and FN values are counted for all classes, but instead of summing these values, precision and recall are calculated for each class. These precision and recall values are then averaged over all classes, optionally weighting each class by its true count of instances. Macro-averaging is appropriate when all classes are equally important. It is also appropriate for an imbalanced data set and ensures that all classes contribute equally. However, it may mask poor performance of important minority classes, and it may lower overall performance measures due to low classifier performance on small or unimportant classes that nonetheless contribute equally to larger or important classes.

Hands-On Exercise

For the multi-class confusion matrix above,

1. Compute precision and recall for each class.
2. Compute the micro-averages of precision and recall and show that they equal the accuracy.
3. Compute the macro-averages of precision and recall.

While precision, recall, and accuracy are useful metrics for evaluating a binary or multinomial classifier, they do not lend themselves to be used directly as loss functions in fitting a model. This is because the assignment of an observation to a class is a function of the probability of its class membership and the actual class assignment may depend on the chosen threshold probability or other decision rule. Thus, a desirable loss function is a function that expresses how close or how different the predicted class membership probabilities are from the true class membership probabilities.

There are two commonly used metrics that quantify such a difference. Both have their origin in information theory and the two are closely related to each other. First, the *cross-entropy* of two probability distributions p_i and q_i over the same set of classes i is defined as:

$$H(p, q) = - \sum_i p_i \log q_i \quad \text{Cross-entropy}$$

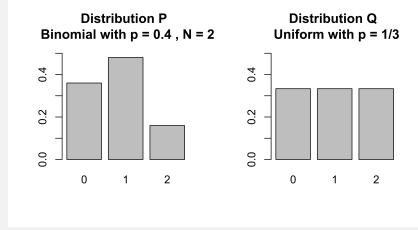
Here, p_i is the true probability of belonging to class i whereas q_i is the predicted probability of belonging to class i . The cross-entropy captures the similarity or difference of the two probability distributions.

The second metric is the *Kullback-Leibler (KL) divergence*. It is effectively the cross-entropy minus the entropy of the true probability distribution, as the following derivation shows:

$$\begin{aligned}
 D_{KL}(P||Q) &= \sum_i p_i \log \left(\frac{p_i}{q_i} \right) && \text{Kullback-Leibler (KL) divergence} \\
 &= \sum_i p_i \log p_i - \sum_i p_i \log q_i \\
 &= -H(p, p) + H(p, q)
 \end{aligned}$$

Hands-On Exercise

1. Consider the two probability distributions P and Q in the following diagrams.



https://commons.wikimedia.org/wiki/File:Kullback-Leibler_distributions_example_1.svg

- (a) Calculate the cross-entropy of P and Q .
 - (b) Calculate the entropy of P .
 - (c) Calculate the KL divergence of P and Q .
- Tip:* Binomial distribution: $\Pr(P = k) = \frac{n!}{k!(n-k)!} p^k (1-p)^{n-k}$
2. Calculate the cross-entropy and KL-divergence for the multi-class confusion matrix above.
 3. Given two probability distributions P and Q over a discrete set of events, where $P = [0.1, 0.4, 0.5]$ and $Q = [0.2, 0.3, 0.5]$, calculate the cross-entropy $H(P, Q)$ and the KL-divergence $D_{KL}(P||Q)$.
 4. In a binary classification task, you have the following probability distributions for the actual labels (P) and predicted labels (Q):
 - $P = [1, 0]$ (the actual class is positive)
 - $Q = [0.7, 0.3]$ (the model predicts a 70% chance of being positive)
 Calculate the cross-entropy loss for this scenario.
 5. Calculate the KL divergence between the following probability distributions:
 - $P = [0.1, 0.9]$
 - $Q = [0.5, 0.5]$

9.6 Resampling Methods

Recall that the goal in prediction is to have an unbiased assessment of the true classification or regression error, and to generalize to prediction of future, yet unseen observations. Finding a suitable prediction model involves two separate steps, that of *model selection*, and that of *model assessment*. Model selection estimates the predictive performance, that is, the error or loss, of different models in order to choose the best one. After having chosen the final model, its prediction error must again be estimated on new data. This is to avoid model selection capitalizing on specific idiosyncrasies of the test data, which may not hold for new, as yet unseen data.

The *validation set approach*, or "holdout" method, then requires not only a training and test data set, but a third set, the validation set. The training data is used to train each of a set of candidate models, the validation data is used to test each candidate model, and the test data is used to evaluate the finally selected model. Typically, the data set is randomly split into 50% training data, 25% validation data, and 25% test data, but other splits are used as well.

One potential problem with this approach is that the validation error can be highly variable and depends critically on the way the data is split. One random split can show very different characteristics than another random split. A second potential problem is that the validation error may overestimate the actual error, because the training set is small, only half of the full data set. Figure 9.12 illustrates the first problem. A single split of the data results in the validation error shown in the left panel. However, repeating the random splitting ten times results in ten different validation errors that may be very different from each other, as shown in the right panel of that figure.

One way to deal with both problems is to use *leave-one-out cross-validation* (LOOCV). In LOOCV, each observation is designated as test observation in turn, while the remaining $n - 1$ observations form the training data set. The model is trained on the training data and tested on the single test observation. This procedure is repeated n times as each observation becomes the test observation in turn. The cross-validation error is

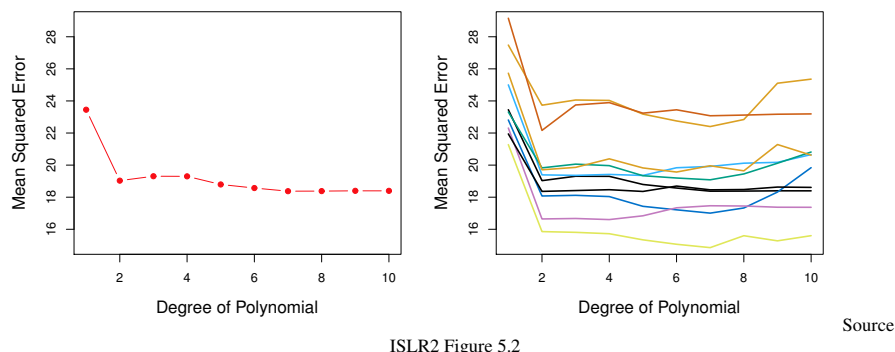


Figure 9.12: Validation error for different random splits of a data set

Source:

ISLR2 Figure 5.2

simply the mean of the n training errors:

$$CV = \frac{1}{n} \sum_{i=1}^n \text{Err}_i$$

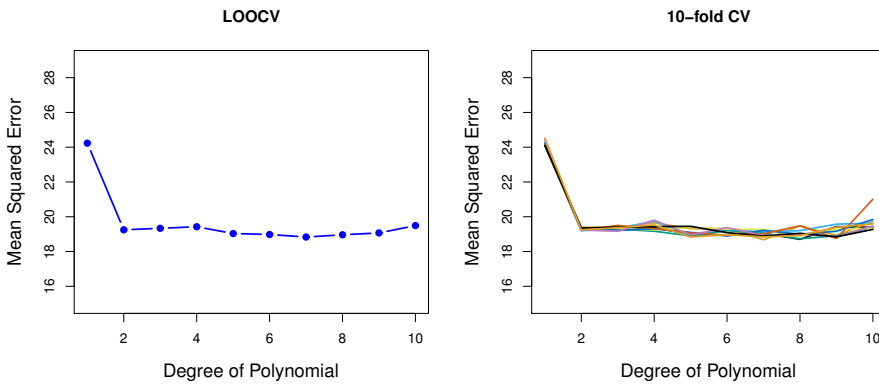
The advantage to LOOCV is that it addresses the two potential problems with the holdout sample approach. Additionally, it is deterministic as there is no randomness to the resulting validation error estimate, because there are no random splits of the data set. Finally, LOOCV shows less overestimation of the validation error rate than the holdout approach. However, a significant drawback is that this approach is computationally expensive, because the model must be fit or trained n times to n different training data sets.

As a compromise between the holdout sample approach and LOOCV, business analysts often use k-fold cross-validation. In this approach, the data set is randomly divided into k sub-samples ("folds"). Each fold is selected as the test data set in turn, with the remaining $k - 1$ folds combining to form the training data set. The cross-validation error is simply the mean of the k cross-validation errors for each of the k test folds. Typically, k is chosen as 5 or 10.

$$CV = \frac{1}{k} \sum_{i=1}^k \text{Err}_i$$

k-fold cross-validation is computationally less expensive and less stable than LOOCV, but it is more stable than the holdout sample approach. The k-fold cross-validation error estimate has a higher bias but lower variance than that of LOOCV. Figure 9.13 illustrates the stability. The left panel shows the LOOCV error estimate for a regression number with different degrees of polynomials. The right panel shows ten different runs each of 10-fold cross-validation (i.e. the model was trained a total of 100 times). The 10-fold cross-validation errors show a much smaller variation than the different holdout sample errors in Figure 9.12.

One important consideration in splitting the data is preventing *information leakage* from training to test or validation data set, in order to ensure that the test and validation data sets are truly independent of the training data. One way in which information could be leaked is when selection of input variables or predictors is done based on characteristics of the entire data set, for example the variance of a variable or the correlation between variables in the entire data set. This selection affects both the training and the test data, essentially leaking some information from the training data to the test data, and thereby making the test data set not truly independent. Another way in which information can leak is by pre-processing variables, like centering around the mean or scaling them to have unit variance. When this is based on the mean or variance of the full data set, information from the training set leaks to the test or validation set, making those data sets not fully independent. As a general rule, any predictor or feature selection and data pre-processing must be done independently for each training set, *after* the split or splits (in the case of k-fold CV) have been made.



Source: ISLR2 Figure 5.4

Figure 9.13: Cross-validation error with LOOCV and 10-fold cross-validation

9.7 Review Questions

Supervised and unsupervised learning

1. Explain the difference between supervised and unsupervised learning. Provide an example for each.
2. Define regression and classification. Discuss one real-life application for each.

Parametric methods

3. What is the difference between a parametric and a non-parametric machine learning model? Provide examples.
4. What are some of the key metrics used to evaluate the quality of a regression model versus a classification model? Discuss their relevance in real-world applications.

Explanation and prediction

5. Define the terms *explanation* and *prediction*. How do they differ in their core objectives when using statistical models?
6. Explain what is meant by the statement that explanatory models are intended to be isomorphic to causal processes.
7. Discuss why explanation models focus on bias minimization and how this affects the model design and interpretation.
8. Provide examples where an explanation model would be more suitable than a prediction model, and vice versa.
9. What are the implications of using a predictive model that does not represent the true causal relationship but still produces accurate predictions?

Bias and variance

10. Explain the terms *bias* and *variance* in the context of statistical modeling. How do they relate to the goals of prediction?
11. Explain the bias-variance tradeoff with an example. You may use a simple regression model as a reference.
12. What are overfitting and underfitting in the context of machine learning? How can each be detected and mitigated?
13. Describe a scenario where a high-bias model would be more appropriate than a low-bias model.
14. Given the following scenarios, identify whether the model is likely suffering from high bias, high variance, or is well-balanced:
 - A model that performs well on training data but poorly on unseen test data.
 - A simple linear regression model that is unable to capture the complexities of the data, resulting in poor performance on both training and test data.
 - A model that performs equally well on training and test data.
15. Describe techniques to reduce bias in a machine learning model.
16. Given a dataset where the relationship between features and target is non-linear and complex, propose a strategy to improve a model that initially has high bias (e.g., linear regression).
17. List and explain strategies to reduce variance in a machine learning model.
18. Imagine you have a deep learning model that performs exceptionally well on the training data but poorly on the validation data. What steps would you take to address this issue?

Regression evaluation

19. Explain the difference between Mean Squared Error (MSE) and Mean Absolute Error (MAE). Why is MAE considered more robust to outliers?
20. Describe the Huber loss function and discuss its advantages over MSE and MAE.
21. Discuss the significance of using test data to evaluate the quality of a regression model. Why is it not advisable to rely solely on training data for model evaluation?

KNN classification

22. Explain how the KNN algorithm estimates the class of a new observation. Include a discussion on the effect of the choice of k .
23. How does changing the value of k in the KNN classifier affect the bias and variance of the model?

Binary classification evaluation

24. Define the terms "precision" and "recall". Provide a scenario where a high recall is more important than high precision, and vice versa.
25. Describe the following metrics and explain their importance in the evaluation of classification models:

- Precision
 - Recall (Sensitivity)
 - F1 Score
 - Specificity
26. Discuss the importance of the ROC curve and AUC in the evaluation of classification models. How do these metrics help in assessing the performance of a model?
27. Given a scenario where you are developing a classifier for a medical diagnosis application, which metric would you prioritize and why?

Multinomial classification evaluation

28. Explain what is meant by multinomial classification. How does it differ from binary classification?
29. Describe the purpose and structure of a confusion matrix in the context of multinomial classification.
30. How is the overall accuracy calculated using a confusion matrix for multinomial classification? Explain using an example.
31. Discuss the difference between micro-averaging and macro-averaging in the context of evaluating classification models.

Cross-entropy and KL-divergence

32. Define cross-entropy and explain its significance in machine learning, especially in classification tasks.
33. Define Kullback-Leibler divergence and explain its relationship with cross-entropy.

Cross-validation

34. Explain the concept of cross-validation and how it helps in model assessment.
35. What is the validation set approach in model evaluation? Describe its potential drawbacks.
36. How does the validation set approach help mitigate the risk of model overfitting?
37. Describe the leave-one-out cross-validation (LOOCV) method. What are the benefits and limitations of using LOOCV for model validation?
38. Compare and contrast LOOCV with the traditional holdout method. In what scenarios might LOOCV be particularly beneficial?
39. Explain k-fold cross-validation. How does it differ from LOOCV in terms of error estimation?
40. Discuss the impact of the number of folds in k-fold cross-validation on the bias and variance of the model error estimate.
41. Define information leakage in the context of data splitting for model evaluation. Why is it important to prevent it?
42. Provide guidelines or methods to prevent information leakage during the preprocessing and feature selection stages of model development.

Chapter 10

Regression and Classification Models

Sources and Further Reading

The material in this chapter is based on the following sources. They are freely available. Consult them for additional information.

Gareth James, Daniel Witten, Trevor Hastie and Robert Tibshirani: *An Introduction to Statistical Learning with Applications in R*. 2nd edition, corrected printing, June 2023. (ISLR2)

<https://www.statlearning.com>

Chapters 2, 3, 4, 5

The book by James et al. provides an easy introduction to machine learning at the introductory undergraduate level. It focuses on applications, not mathematics, and contains many exercises using R. Concepts are well explained and illustrated. There is a similar book available by the same authors with applications in Python. This book is a more accessible of the following book.

Trevor Hastie, Robert Tibshirani, and Jerome Friedman: *The Elements of Statistical Learning*. 2nd edition, 12th corrected printing, 2017. (ESL)

<https://hastie.su.domains/ElemStatLearn/>

Chapters 2, 3, 4, 7

The book by Hastie et al. still sets the standard for statistical learning. It is widely used and cited. Its treatment is more technical than the previous book and there are

no exercises in R or Python. However, it covers the concepts in more depth (and a few more formulas). However, it is still very accessible even to an undergraduate audience.

Kevin P. Murphy: *Probabilistic Machine Learning – An Introduction*. MIT Press 2022.

<https://probml.github.io/pml-book/book1.html>

Chapters 4, 6, 9, 10, 11

Murphy's book is available under a creative-commons license. It is a somewhat more technical treatment of the material, but with many illustrations and examples. It is quite comprehensive in its coverage and targeted at the advanced undergraduate or graduate student.

10.1 Introduction

This chapter is an introduction to supervised machine learning using R and includes both regression and classification problems. The chapter introduces linear regression and penalized regression models (ridge regression and LASSO). For classification, it introduces the methods of logistic regression and k-Nearest-Neighbours, which was already briefly discussed in the previous chapter. This chapter builds on the introductory material to supervised learning in the previous chapter.

10.2 Linear Regression

Linear regression fits a simple statistical model to a set of input and output variables. The true model is assumed to take the form:

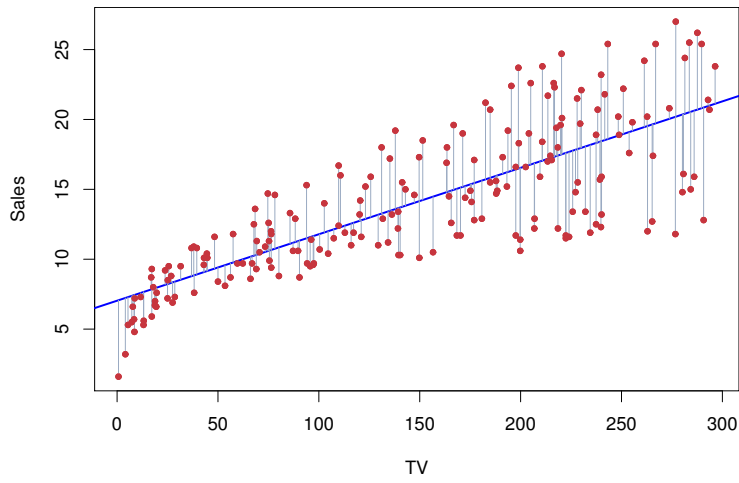
$$f(X) = Y = \beta_0 + \beta_1 X + \epsilon$$

while the fitted, approximate model is:

$$\hat{f}(X) = \hat{Y} = \hat{\beta}_0 + \hat{\beta}_1 X$$

The values of $\hat{f}(X) = \hat{Y}$ are called the *fitted values* in statistics or the *predicted values* in machine learning. The difference in terminology reflects the fact that traditional statistics looks back at the training data for the output values determined by the model after it has been fit to the training data, whereas machine learning emphasizes prediction of output values for new inputs.

This equation shows the form of the assumed functional relationship between X and Y is linear in the parameters β . In other words, there are no polynomials or other transformations of β , such as β_1^2 or $\log \beta_1$. It is the linearity of the parameters that



Source: ISLR2 Figure 3.1

Figure 10.1: A linear regression model

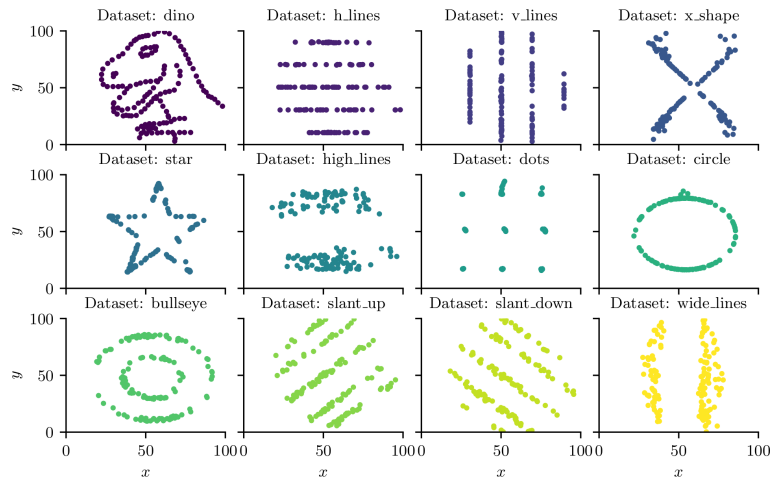
makes a regression linear, *not* the linearity of the predictors. For example, adding a polynomial term $\beta_2 X^2$ to the right-hand side of this regression equation would still make this a linear regression.

Figure 10.1 shows the fitted regression line expressed by the intercept ($\hat{\beta}_0$) and slope ($\hat{\beta}_1$) parameters. The distance between this line and the data points (coloured red) represents the error term ϵ in the regression equation. Importantly, the error is the vertical differences in the Y direction between fitted regression line and the data point, *not* the shortest distance between the regression line and the data point.

The first task in performing a regression analysis is to identify the form of the regression model or regression equation. While Figure 10.1 fitted a model that is linear in X , it is clear that this model is not a good fit for small X values, as the corresponding observations are all below the regression line.

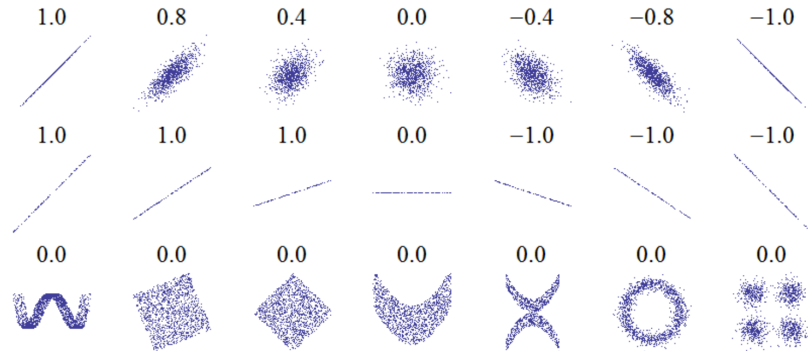
To identify an appropriate functional form, it is insufficient to simply examine summary statistics like correlations or covariances between variables. Figure 10.2 shows twelve data sets with the same correlation between the two variables. It is clear that none of them can be fitted to a simple moodel that is linear in X . While some, like the circle or the bullseye data set, can be appropriately transformed and fitted to linear regression models that involve polynomials or other functions of X , it is not clear what functional form the dinosaur head might take.

It is also insufficient to fit a model that is linear in X and use the fit as indication for the appropriateness of the model. Consider the data sets shown in Figure 10.3. The correlations between the two variables are shown above each data set. Visual inspection is sufficient to show that data sets with the same correlation do not necessarily have the same regression slope, or should even be fitted to the same linear model.



Source: Murphy Figure 2.6

Figure 10.2: The "Datasaurus Dozen" – All datasets have the same correlation between the two variables



Source: Murphy Figure 3.1

Figure 10.3: Datasets with the same correlation (as indicated above each dataset) between two variables do not need to have the same regression slope

The objective for estimating the parameters β_0 and β_1 from the data set is to minimize the residual or error, that is, the difference between f and \hat{f} . So that positive and negative differences do not compensate for each other, the difference is either squared or the absolute value is taken. The former is called *mean squared error* (MSE) while the latter is called *mean absolute error* (MAE). In statistics, a closely related concept is the *residual sum of squares* (RSS). The MSE is the RSS divided by the number of observations n , and minimizing one also minimizes the other. While the discussion in

the previous chapter indicated that the MAE is more robust in that outliers, i.e. observations with large errors, have less influence on the estimated values of the parameters, it is customary to fit linear regression models using the RSS or the MSE.

$$\begin{aligned} RSS &= \sum_i (y_i - \hat{\beta}_0 - \hat{\beta}_1 x_i)^2 \\ MSE &= \frac{1}{n} RSS \end{aligned}$$

The linear regression model is simple enough that an optimal solution can be derived analytically. The optimal least-squares estimates are:

$$\begin{aligned} \hat{\beta}_1 &= \frac{\sum_i (x_i - \bar{x})(y_i - \bar{y})}{\sum_i (x_i - \bar{x})^2} \\ \hat{\beta}_0 &= \bar{y} - \hat{\beta}_1 \bar{x} \end{aligned}$$

where \bar{x} and \bar{y} are the sample means.

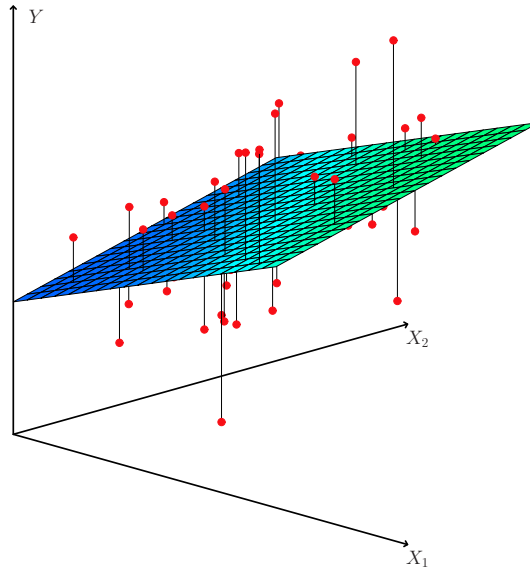
The parameter value estimates have *standard errors* associated with them that indicate the uncertainty of these estimates. This is based on the idea that the training data set is a small random sample from the overall population, and taking other random samples may well yield slightly different parameter values.

Using the standard errors of the estimates, a *t statistic* can be calculated from the difference of the estimate of a parameter $\hat{\beta}$ and some value V .

$$t = \frac{\hat{\beta} - V}{SE(\hat{\beta})}$$

The statistic t is a random variable whose values are distributed according to a student-t probability distribution. This allows one to calculate the probability of observing a value of t or larger under the assumption that $\hat{\beta} = V$, that is, there is no difference between V and the estimate of the parameter $\hat{\beta}$. If the calculated probability is very small, it is unlikely that this assumption holds and one might conclude that $\hat{\beta} \neq V$. This procedure is called the *t-test*. In most applications of this test to regression model parameters, V is set to 0. Because for large sample sizes the student-t distribution approaches the normal (Gaussian) probability distribution, the probability is sometimes calculated from this distribution, the statistic is called the *z statistic* and the test is then called a *z-test*.

Another important statistic in linear regression analysis is the proportion of explained variance, designated as R^2 . It is defined as:



Source: ISLR2 Figure 3.4

Figure 10.4: Example linear regression with two predictors

$$R^2 = \frac{TSS - RSS}{TSS} = 1 - \frac{RSS}{TSS}$$

where $TSS = \sum_i(y_i - \bar{y})^2$ is the total sum of squares. An R^2 value close to 1 indicates that the fitted regression model explains a large proportion of the variability in the data set, that is, it explains the data well. In contrast, a value close to 0 indicates that the fitted regression model explains very little of the observed variability, it does not explain the observed data well. Another important interpretation of the R^2 is as the correlation between the true Y and the fitted or predicted values \hat{Y} .

Adding additional predictors into the linear regression model is straightforward, as is the inclusion of qualitative or categorical predictors. For example, a model with two predictors X_1 and X_2 assumes the true form

$$Y = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \epsilon$$

and fits a plane to a set of points in three dimensional (X_1, X_2, Y) space, as shown in Figure 10.4.

Qualitative predictors (called *factors* in statistics) with multiple, exclusive *levels* or categories can be included using *dummy variables*. A categorical variable that can take

on k values requires $k - 1$ binary dummy variables. For example, a factor x that has four levels "a", "b", "c", and "d" might be encoded using three dummy variables as follows:

$$\begin{aligned}x_{i1} &= \begin{cases} 1 & \text{level "a"} \\ 0 & \text{else} \end{cases} \\x_{i2} &= \begin{cases} 1 & \text{level "b"} \\ 0 & \text{else} \end{cases} \\x_{i3} &= \begin{cases} 1 & \text{level "c"} \\ 0 & \text{else} \end{cases}\end{aligned}$$

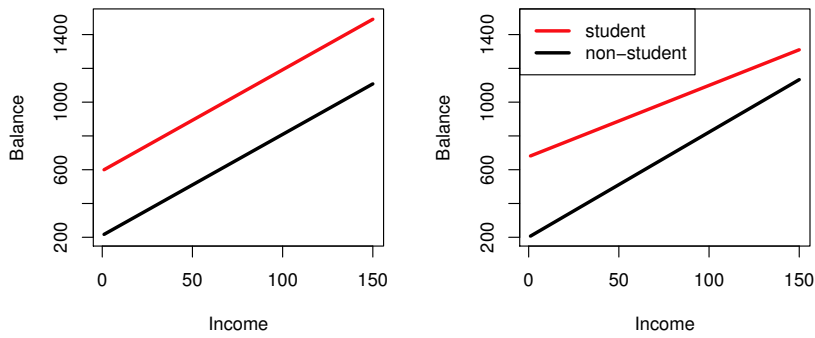
Note that $x_{i1} = x_{i2} = x_{i3} = 0$ represents level "d". Such *contrasts* determine how factor levels are coded using dummy variables.

Earlier, it was noted that a linear model is linear in its parameters, not necessarily in its input variables. Hence, it is possible to include transformations of input variables in a linear regression model, such as polynomials, products, or other functional transformations like logarithms, square roots, etc. For example, the model

$$Y = \beta_0 + \beta_1 X_1 + \beta_2 X_1^2 + \beta_3 X_2 + \beta_4 X_1 X_2 + \epsilon$$

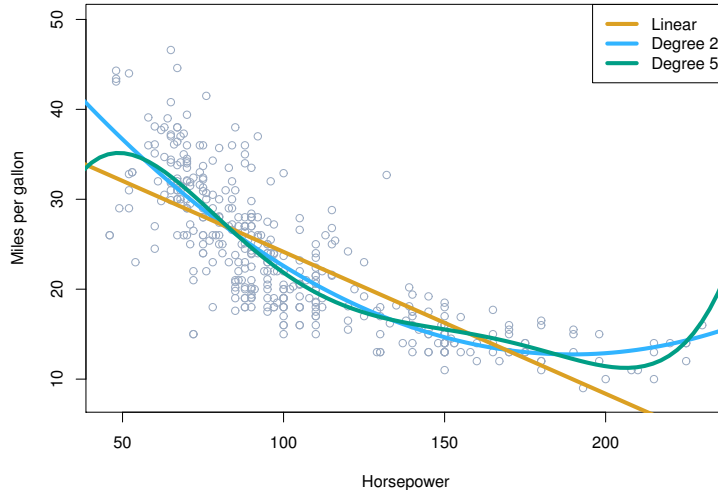
is still linear in its parameters β_i . This model also shows the difference between *input variables* and *predictors* or *features* that was alluded to in the previous chapter. This model has two input variables X_1 and X_2 but has four predictors or features, X_1 , X_1^2 , X_2 , and $X_1 X_2$. In statistics terminology, the coefficients of single input variables, here β_1 and β_3 , are said to represent *main effects*, while the parameters of products of two or more input variables, here β_4 are said to represent *interaction effects*. Interaction effects can include the product of a numerical variable and a dummy binary variable. This results in different regression slopes for different categories or factor levels, as shown in the example in Figure 10.5 where the left panel shows no interaction effect, that is, the lines are parallel with the same slopes. The right panel shows an interaction effect where the slopes for different categories are different.

An example of polynomial predictors like X^2 , X^3 or higher degrees is shown in Figure 10.6. As the figure shows, and as is true in general, increasing the number of predictors increases the flexibility of the model to fit the data. This results in smaller model bias but at the expense of model variance.



Source: ISLR2 Figure 3.7

Figure 10.5: Example interaction effect in linear regression



Source: ISLR2 Figure 3.8

Figure 10.6: Regression example with polynomial predictors

10.3 Linear Regression in R

Linear regression is part of the basic R system, no libraries need to be installed. The function `lm()` requires a formula representing the regression model and a data frame, and returns a linear model.

The following example uses the `Boston` data set that contains housing prices ("median value", `medv`) as well as demographic and socioeconomic information for suburbs in the city of Boston. The data is available as part of the `ISLR2` package¹.

¹The R code for this example is based on material in Section 3.6 of ISLR2

First, examine the data, get summary statistics, examine the first few rows of the data frame, and create scatterplots to identify the form of a linear model:

```
# Data set from the textbook 'Introduction to
# Statistical Learning with Applications in R'
library(ISLR2)

# Get a description of the data
?Boston

# Get a summary and examine first few rows
summary(Boston)
head(Boston)

# Bivariate scatterplots
plot(Boston)
```

The formula interface to `lm()` is the easiest to use and resembles the way one would write the regression equation. The `~` sign separates the output on the left side from the predictors or features on the right side of the formula. A `1` in the formula represents the intercept (β_0 in the formula above). The intercept is normally added automatically but can be explicitly added or removed (using `-1`) as needed.

The following R code fits a simple model to predict the median house value, shows the model summary, produces some plots, and illustrates the use of the `predict()` function to predict values:

```
# Fit a model with intercept only
fitted.model <- lm(medv ~ 1, data=Boston)
summary(fitted.model)

# Fit a model with predictor lstat
fitted.model <- lm(medv ~ lstat, data=Boston)
summary(fitted.model)

# Plot the data and the regression line
plot(medv ~ lstat, data=Boston)
abline(fitted.model, lwd=3, col='red')

# Plot the residuals against predicted values
plot(predict(fitted.model), residuals(fitted.model))

# Predict three new observations of lstat
predict(fitted.model, data.frame(lstat=c(5, 10, 15)),
        interval='confidence')
```

The following R code fragment adds a second input variable (`age`) to the model. It also demonstrates some special notation in the R formula interface, such as the `.` to include all main effects, the `:` to specify interaction effects, and the `*` to include both main and interaction effects.

```

# Add another predictor
fitted.model <- lm(medv ~ lstat + age, data=Boston)

# Add all main effects
fitted.model <- lm(medv ~ ., data=Boston)

# Add interaction terms
fitted.model <- lm(medv ~ lstat + age + lstat:age, data=Boston)

# Shorter and equivalent
fitted.model <- lm(medv ~ lstat*age, data=Boston)
summary(fitted.model)

```

The next example adds polynomial terms to the regression. It uses the `I(.)` function in R that can also be used with other transformations of the inputs, such as `I(sqrt(.))` or `I(log(.))`. To make it easier to include all polynomials up to a particular degree, one can us the `poly(.)` function:

```

# Add a polynomial term; use the I(.) function
# for any data transformations, such as log(),
# or exp() or sqrt() as well as polynomials
fitted.model <- lm(medv ~ lstat + I(lstat^2), data=Boston)
summary(fitted.model)

# Add all polynomial terms up to degree 5
fitted.model <- lm(medv ~ poly(lstat, 5), data=Boston)
# Note the coefficients for the polynomials in the summary
summary(fitted.model)

```

The use of categorical input variables is illustrated in the following example that uses the `Carseats` data, also from the `ISLR2` package. The data frame represents sales data of car seats for different stores and a range of store characteristics in other variables, many of which are categorical.

The following R code demonstrates functions for dealing with categorical variables, called "factors" in R. The example then fits a regression model to predict Sales from the the main effects of all input variables in the data frame.

```
?Carseats

# Identify factor/categorical variables and their levels:
is.factor(Carseats$ShelveLoc)
levels(Carseats$ShelveLoc)
levels(Carseats$Urban)
levels(Carseats$US)

# Contrasts show the dummy variables created (columns) and
# the values they take for different factor levels (row)
contrasts(Carseats$ShelveLoc)
contrasts(Carseats$US)

# Fit the model
summary(lm(Sales ~ . , data=Carseats))
```

Hands-On Exercise

Use the `Auto` data set from the `ISLR2` library with `mpg` as the target.

1. Perform a linear regression with `horsepower` as predictor
2. Is there a relationship between the predictor and target? What form and how strong?
3. What is the predicted `mpg` value for a `horsepower` of 98?
4. Plot the response and predictor. Use the `abline()` function to add the regression line
5. Produce a scatterplot of all variables
6. Perform a linear regression of all main effects (except for the variable name), then remove non-significant predictors
7. Use the `*` and `:` symbols to add interaction effects. Retain only significant ones
8. Add transformations of the predictors (using the `I(.)` function) such as $\log(X)$, \sqrt{X} , X^2 .

Source: ISLR2 Section 3.7

Hands-On Exercise

Use the `Carseats` data set from the `ISLR2` library with `Sales` as the target.

1. Perform a linear regression with `Price`, `Urban` and `US` as predictors
2. Interpret the coefficients. Tip: Some variables are categorical
3. Remove non-significant predictors
4. How well do the two models fit the data?
5. Determine the 95% confidence intervals for the coefficients of each model.
6. (How) does the importance of predictors change?

Source: ISLR2 Section 3.7

10.4 Cross-Validation in R

The previous chapter already introduced the concepts of cross-validation. Recall that the appropriate way to judge the predictive performance of a model is not to evaluate it on its training data, but to evaluate it on unseen test data. This section illustrates three different cross-validation approaches in R, beginning with the validation set or holdout sample approach, which splits the data set randomly into training and test data.

The following example uses the `Auto` data set from the `ISLR2` package, which contains information on different vehicle models, their fuel economy ("miles per gallon", `mpg`) and vehicle characteristics².

Randomly splitting the data set is accomplished using the `sample()` function in R which returns a boolean vector with values `TRUE` or `FALSE` that can be used to select the appropriate data from the data frame.

To make the example repeatable, the initial seed of the pseudo-random number generator (RNG) is set to a fixed value with `set.seed()`. Computers cannot generate truly random numbers; they are deterministic machines. The pseudo-random numbers they generate are computed by a deterministic algorithm, the "random number generator" (RNG), based on a given start value. With the same start value, the algorithm produces the same sequence of numbers. A good RNG produces numbers that are effectively indistinguishable from true random numbers.

²The R code for this example is based on material in Section 5.3 of ISLR2

```

# Set the seed for the pseudo-random
# number generator (RNG)
set.seed(1)

# Randomly use half the Auto data as training sample
train.idx <- sample(nrow(Auto), nrow(Auto)/2)
train.data <- Auto[train.idx,]
test.data <- Auto[-train.idx,]

# Fit model to (train model on) a subset
fitted.model <- lm(mpg ~ horsepower, data=train.data)

# Calculate the test data MSE by predicting from the test data set
mean((test.data$mpg-predict(fitted.model,test.data))^2)

# Calculate the training MSE by predicting from the training data set
mean((train.data$mpg-predict(fitted.model,train.data))^2)

# The MSE can also be calculated from the squared residuals
mean(summary(fitted.model)$residuals^2)

```

The next R code block illustrates Leave-One-Out Cross-Validation (LOOCV), where one observation is used as test observation, and the remainder form the training data. This is repeated so that each observation becomes the test observation in turn. The errors are then averaged.

While this can be done manually in R using an iteration over the data frame, the library `boot` provides easy-to-use cross-validation functions for models fitted using the `glm` function (generalized linear models). Note that LOOCV is just k-fold cross-validation where k , the number of test data folds, is equal to the number of observations.

```

library(boot)

# Fit a model with glm and show its summary
fitted.model <- glm(mpg ~ horsepower, data=Auto)
summary(fitted.model)

# LOOCV is k-fold CV where k equals the number of observations
cv.err <- cv.glm(Auto, fitted.model, K=nrow(Auto))
cv.err$delta[1]

```

The same functions can be used for k-fold CV with a typical value for k :

```

cv.err <- cv.glm(Auto, glm.fit, K=10)
cv.err$delta[1]

```

Cross-validation is useful for comparing different models, as shown in the following example that fits linear regression models with different degrees of polynomials to a data frame and computes their cross-validation errors using a `for` loop in R:

```

set.seed(1)
cv.err <- rep(0, 5)
for(i in 1:10) {
  fitted.model <- glm(mpg ~ poly(horsepower,i), data=Auto)
  cv.err[i] <- cv.glm(Auto, fitted.model, K=10)$delta[1]
}
print(cv.err)

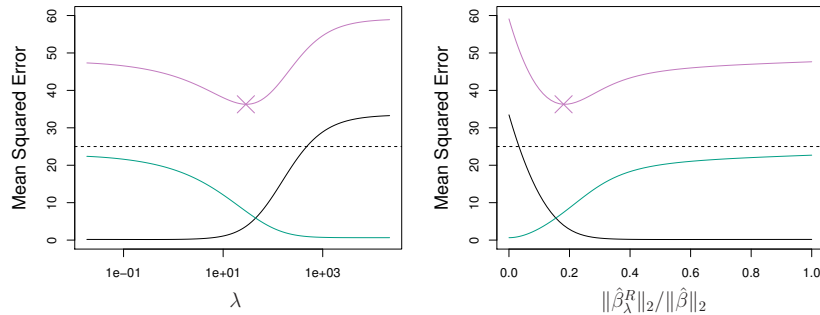
```

Hands-On Exercise

1. Fit a regression model to the Boston data set with `medv` as target, and `age`, `lstat`, and `ptratio` as predictors
2. Using the holdout approach, compute the test error of this model. Perform the following steps
 - (a) Split the data set using 75% for training and 25% for testing
 - (b) Fit the model to training data
 - (c) Predict the target for the testing data
 - (d) Compute the test error
3. Repeat the previous step 3 times, using different splits. How do the results change?
4. Calculate the mean and the variance of the test errors of the four splits.
5. Include `dis` as predictor in the model. Does it reduce the test error?
6. Calculate the test error estimate using LOOCV. Compare your result to the mean that you computed in step 4.
7. Calculate the test error estimate using 4-fold cross-validation. Compare the estimate to the mean that you computed in step 4

10.5 Shrinkage Methods

Shrinkage methods are so-called because their aim is shrink the magnitude of the regression parameter values $\hat{\beta}_i$. This is primarily to avoid overfitting a model to one specific data set. In other words, they are a kind of *regularization*. Shrinkage methods for linear regression model works by *penalizing* the model for large regression parameter values, that is, by performing *penalized regression*. There are three frequently used types of penalized regression or linear regression regularization: L1 regularization penalizes large absolute parameter values (that is, the L1 norm of the vector of parameters $\hat{\beta}$), L2 regularization penalizes large squared parameter values (that is, the L2 norm of the vector of parameters $\hat{\beta}$), and Elastic Net regularization is a combination of both. L1 regularization is also known as the *LASSO* ("least absolute shrinkage and selection operator") while L2 regularization is known as *ridge regression* or Tikhonov regularization.



Source: ISLR2 Figure 6.5

Figure 10.7: Bias (black), variance (turquoise), and MSE (pink) in ridge regression

10.5.1 Ridge Regression

The loss function or minimization objective of ridge regression is the usual RSS that now also includes a term that penalizes large values of β .

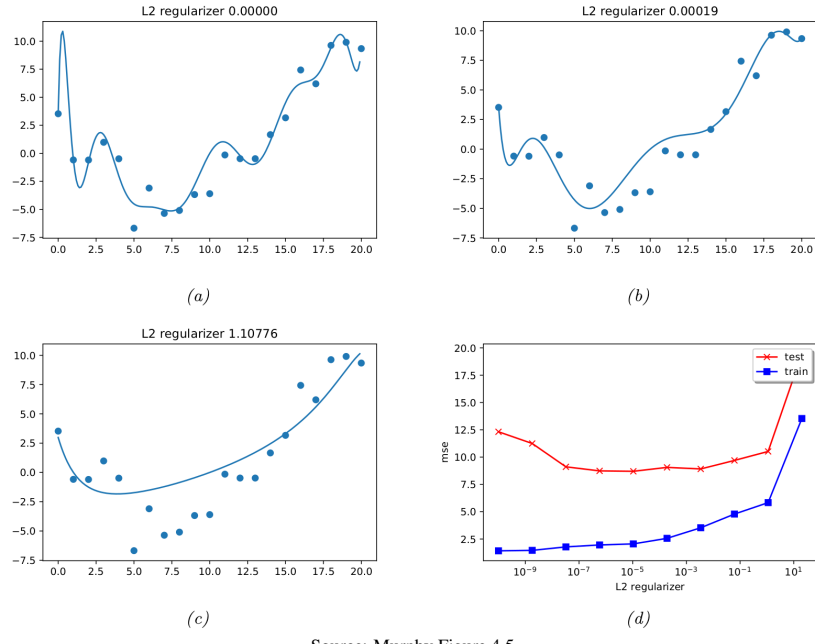
$$\text{Minimize } RSS + \lambda \sum_{j=1}^p \beta_j^2 = RSS + \lambda \|\beta\|_2^2$$

where $\|\cdot\|_2$ indicates the *L2 vector norm*, that is, the sum of squared entries of the vector β .

Because the scale, that is, the standard deviation or variance, of different predictors affects the size of their associated β parameter values, *all predictors should be rescaled or standardized to have the same standard deviation prior to performing a ridge regression.*

The degree of penalization or the amount of shrinkage is controlled by the parameter λ . Larger values for λ increase the penalty, thus generally leading to larger model bias but smaller variance. This effect is shown in Figure 10.7 where the left panel shows the bias (black line), the variance (turquoise line) and the total MSE (pink line) as a function of λ and the right panel shows the bias, the variance, and the total MSE as a function of the proportion to which the L2 norm of the β is restricted or penalized compared to the unrestricted estimates, i.e. $\|\hat{\beta}_\lambda^R\|_2 / \|\hat{\beta}\|_2$ where $\|\cdot\|_2$ indicates the *L2 vector norm*, i.e. the sum of the squared elements of the vector.

Figure 10.8 shows this effect for a polynomial of degree 14 that is fitted to data set of 21 observations. The different panels show different degrees of penalization or shrinkage as indicated by different values of λ . The bottom right panel in Figure 10.8 shows the MSE on the training data (blue) and test data (red) for multiple values of λ . It is clear that the unpenalized model in panel (a) of Figure 10.8 overfits, while models that are



Source: Murphy Figure 4.5

Figure 10.8: Fitting a degree 14 polynomial with ridge regression

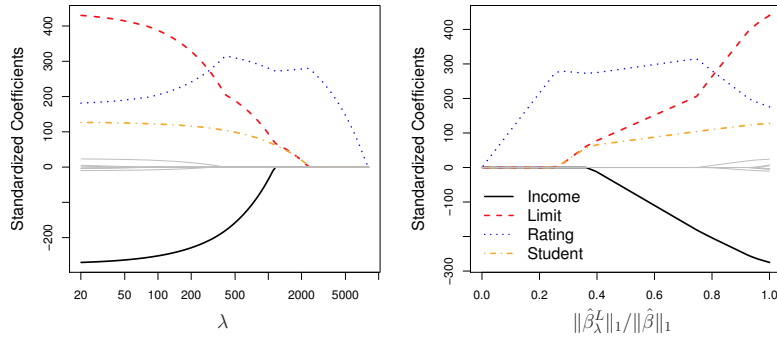
penalized too heavily underfit and have large bias, such as the model in panel (c) of Figure 10.8.

10.5.2 LASSO

In contrast to the ridge regression, where the model parameter values are shrunk towards zero, but are never forced to equal zero, the LASSO does just that. In this form of penalized regression, as the degree of penalty is increased, model parameter values are set to 0, effectively making the LASSO a model or *predictor selection method* as well, that is, it can be used to select only important predictors. This is reflected in its name: "Least Absolute Shrinkage and Selection Operator". The advantage over ridge regression is that this results in more parsminious, that is, smaller, models that are easier to interpret.

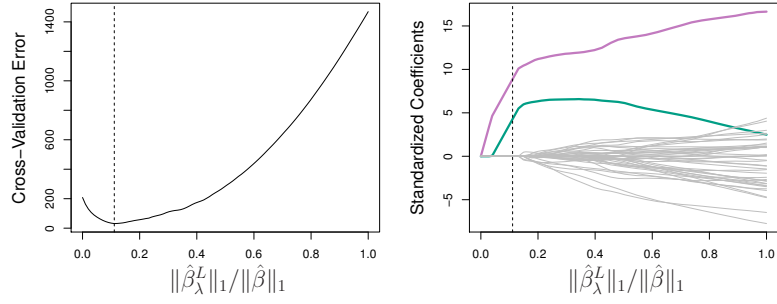
The loss function or minimization objective of the LASSO is the RSS but penalized for the L1 norm of the vector of parameters β . As in ridge regression, the amount of shrinkage is controlled by a parameter λ .

$$\text{Minimize } RSS + \lambda \sum_{j=1}^p |\beta_j| = RSS + \lambda \|\beta\|_1$$



Source: ISLR2 Figure 6.6

Figure 10.9: Predictor selection in the LASSO



Source: ISLR2 Figure 6.13

Figure 10.10: Cross-validation error in the LASSO

where $\|\cdot\|_1$ indicates the *L1 vector norm*, that is the sum of the absolute values of the elements of the vector β .

The model selection property of the LASSO are shown in Figure reffig:lasso1. The left panel shows the value of different model parameters as a function of the penalization parameter λ . As λ increases, fewer and fewer parameters are allowed to retain absolute values larger than 0. The same is shown in the right panel of Figure 10.9, but now as a function of the relative size of the restricted L1 norm of β compared to the L1 norm of the unconstrained β .

The LASSO and ridge regression shrinkage or penalty parameter λ is typically chosen through cross-validation to minimize the cross-validation or test error. For cross-validation, a "grid" or range of possible values of λ is defined. A model is fitted for each value of λ and its cross-validation or test error is calculated. The final model is then fitted using the optimal value of λ , the is, the value that results in the lowest cross-validation or test error.

This is illustrated in Figure 10.10 for a LASSO, where the left panel shows the cross-

validation error as a function of the relative shrinkage. Towards the left of the graph in that panel, where shrinkage is maximal, the model underfits, resulting in a relatively large error due to a large bias, whereas to the right of the graph, where shrinkage is minimal, the model overfits, leading to a large cross-validation error due to large variance. The right panel of Figure 10.10 shows the size of model parameter values; at the optimal λ only two model parameters are different from zero, that is, only two predictors are selected for the model.

10.5.3 Elastic Net

The Elastic Net is a combination of ridge regression and LASSO, controlled by the parameter α . The Elastic Net penalty is defined as

$$\lambda (\alpha \|\beta\|_1 + (1 - \alpha) \|\beta\|_2^2)$$

When $\alpha = 0$ the Elastic Net reduces to a ridge regression, and for $\alpha = 1$ the Elastic Net reduces to the LASSO. The optimal combination of α and λ is found through cross-validation as described above.

10.6 Shrinkage Methods in R

The `glmnet` library for R implements the Elastic Net which can be used for both ridge regression and LASSO as well, simply by choosing the appropriate value for the parameter α .

The following R code examples use the `Hitters` data set containing information on baseball players. The data set is part of the `ISLR2` library. The code examples model a player's Salary as the output or prediction target and use a number of other variables as inputs³.

```
library(ISLR2)
library(glmnet)

# Remove missing values
Hitters <- na.omit(ISLR2::Hitters)
```

The `glmnet(.)` function requires separate x (predictors) and y (target) values, instead of providing a formula interface like the `lm(.)` and `glm(.)` functions. To create dummy variables for categorical variables, use the `model.matrix` function:

³The R code for this example is based on material in Section 6.5.2 of ISLR2

```
# Create dummy variables for categorical variables
# and remove the intercept (first column) from the model
x <- model.matrix(Salary ~ ., Hitters)[, -1]
y <- Hitters$Salary
```

To illustrate the concepts of ridge regression, the following R code sets up a grid of 100 different λ values, fits ridge regression models to each of them, and shows information about two models for different λ values:

```
grid <- 10^seq(from=-10, to=-2, length=100)
print(grid)
ridge.model <- glmnet(x, y, alpha=0, lambda=grid)

# Select the 50th lambda value
ridge.model$lambda[50]
coef(ridge.model)[, 50]
L2.norm = sqrt(sum(coef(ridge.model)[-1, 50]^2))

# Select the 60th lambda value
ridge.model$lambda[60]
coef(ridge.model)[, 60]
L2.norm = sqrt(sum(coef(ridge.model)[-1, 60]^2))
```

The optimal value for λ is chosen through cross-validation. For this, a holdout test data set is created to evaluate the final model. The training data set portion is used with cross-validation, so that the final model evaluation is done on an independent data set from model selection.

```
# Randomly split the Hitters data
train.idx <- sample(nrow(Hitters), nrow(Hitters)/2)
x.train <- x[train.idx,]
x.test <- x[-train.idx,]
y.train <- y[train.idx]
y.test <- y[-train.idx]
```

The `glmnet` library provides the function `cv.glmnet` as an easy-to-use way to combine Elastic Net model fitting with k-fold cross-validation. The following example uses 5-fold CV on the training portion of the data set:

```
# 5-fold cross-validation, use MSE as metric
cv.out <- cv.glmnet(x.train, y.train, alpha=0,
nfolds=5, type.measure='mse')
```

The next R code block shows the optimal λ and plots the MSE for different values of λ . The generated plot is shown in Figure 10.11. It includes the standard errors of

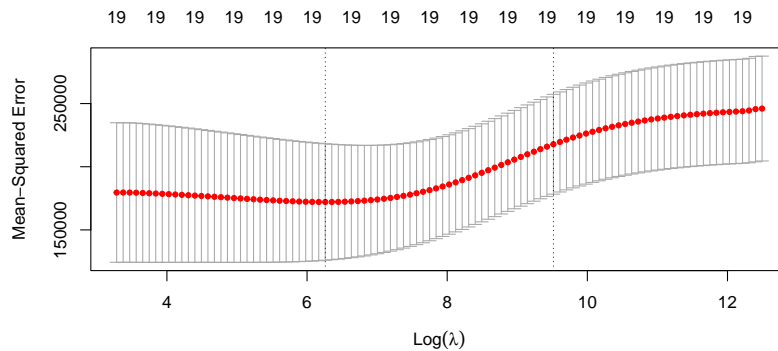


Figure 10.11: Cross-validation MSE in ridge regression

the MSE (as determined by cross-validation) and the number of non-zero coefficients (indicated above the graph). The left vertical line indicates the optimal λ , while the right vertical line indicates that largest λ for which the error is within one standard error of the minimum error, that is, of the error of the optimal λ .

```
print(cv.out)
plot(cv.out)
lambda.opt <- cv.out$lambda.min
```

To evaluate the final model, the holdout test data is fitted to a ridge regression model with the optimal λ . Then, the model parameter values of the ridge regression with the optimal λ are compared to the parameter values of an unrestricted linear regression model. Note the use of the `type='coefficients'` parameter to `predict(.)` that asks for the model parameter values, rather than the predicted target values in the following R code block.

```
# Fit test data:
ridge.test <- glmnet(x.test, y.test, alpha=0)
# Show the coefficients at optimal lambda
predict(ridge.test, type='coefficients', s=lambda.opt)
# Compare to unpenalized least-squares fit
coef(lm.fit(x.test, y.test))
```

Finally, the target values for the holdout test data set are predicted. Note the use of the `type='response'` parameter to `predict(.)` that asks for the predicted target values, rather than the model parameter values, in the following R code block:

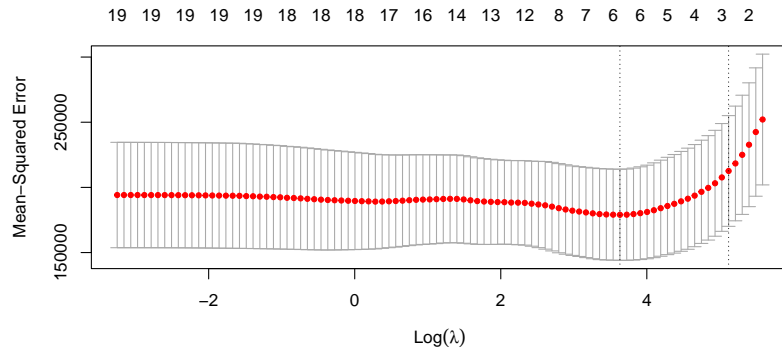


Figure 10.12: Cross-validation MSE in the LASSO

```

predictions <- predict(ridge.test, type='response',
                      s=lambda.opt, newx=x.test)
# Calculate test MSE to compare to the CV optimal MSE above:
mean((predictions - y.test)^2)

```

Because it uses the same `glmnet()` and `cv.glmnet()` functions, the LASSO in R is very similar to ridge regression. The following example shows cross-validation on the training data set to determine the optimal value for λ .

```

# Set alpha to 1 for lasso
cv.out <- cv.glmnet(x.train, y.train, alpha=1,
                     nfolds=5, type.measure='mse')
print(cv.out)
plot(cv.out)
lambda.opt <- cv.out$lambda.min

```

The MSE values for different values of λ are shown in Figure 10.12. Note that now the number of non-zero model parameters, indicated above the graph, decreases. At the optimal λ value, there are only six non-zero parameters, that is, only six predictors are selected to remain in the model.

Hands-On Exercise

Predict the number of applications received using the other variables in the College dataset

1. Split the data set into a training and a test set
2. Fit an unpenalized linear model on the training set. Report the test error.
3. Fit a ridge regression model on the training set, with λ chosen by cross-validation. Report the test error.
4. Fit a lasso model on the training set, with λ chosen by cross-validation. Report the test error.
5. Compare and contrast the results

Source: ISLR2, Section 6.6

Hands-On Exercise

Predict the per-capita crime rate in the Boston data set using the other variables.

1. Split the data set into a training and a test set
2. Fit an unpenalized linear model on the training set. Report the test error.
3. Fit a ridge regression model on the training set, with λ chosen by cross-validation. Report the test error.
4. Fit a lasso model on the training set, with λ chosen by cross-validation. Report the test error.
5. Compare and contrast the results

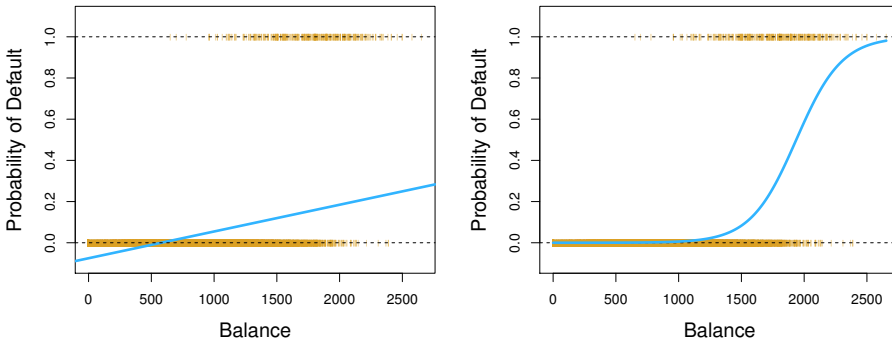
Source: ISLR2, Section 6.6

10.7 Classification

In classification, the output or target value is categorical. In particular, in binary classification, the target may take on one of two values. Classification predicts class membership for new observations by estimating the probability of membership in each class for that observation.

10.7.1 Logistic Regression

Linear models, as used in regression, are not suitable for classification without modification, because probabilities are bounded between 0 and 1, inclusive, while the regression output can take on any real value. The solution to this problem is to use a *link function* that transforms the output of a linear regression model and bounds it between 0 and 1. This is shown in Figure 10.13 where the probability of credit card default are to be predicted from the credit card balance. The yellow points are the training observations, classified as either defaulters (0) or non-defaulters (1). The left panel shows that a linear regression (blue line) yields negative probabilities for small balances, and



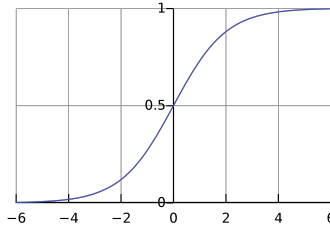
Source: ISLR2 Figure 4.2

Figure 10.13: Transforming linear regression output for binary classification

will yield probabilities larger than one for very large credit card balances. The right panel shows the transformed linear regression output, now bounded between 0 and 1, and interpretable as probabilities from which class membership can be predicted.

There are many transformation or link functions that may be used, but a popular one is the *logistic function*, a type of *sigmoid function* ("s-shaped") (often the two terms are equated), defined as follows:

$$\begin{aligned}\sigma(x) &= \frac{1}{1 + e^{-x}} \\ &= \frac{e^x}{1 + e^x} \\ &= 1 - \sigma(-x)\end{aligned}$$



<https://commons.wikimedia.org/wiki/File:Logistic-curve.svg>

For the binary case, this leads to *binary logistic regression*, expressed by the following equalities:

$$\begin{aligned}p(X) &= \sigma(\beta_0 + \beta_1 X) \\ &= \frac{e^{\beta_0 + \beta_1 X}}{1 + e^{\beta_0 + \beta_1 X}}\end{aligned}\tag{10.1}$$

$$\Rightarrow \frac{p(X)}{1 - p(X)} = e^{\beta_0 + \beta_1 X} \quad \text{"Odds"}\tag{10.2}$$

$$\Rightarrow \log\left(\frac{p(X)}{1 - p(X)}\right) = \beta_0 + \beta_1 X \quad \text{"Log-Odds", "Logits"}\tag{10.3}$$

Equation 10.1 defines the probability of an observation being "true" as just the logistic

transformation of the linear combination of predictors. Dividing and a little algebraic rearrangement yields equation 10.2 which represents the *odds* as the exponential of the linear combination of predictors. Taking the natural logarithm yields equation 10.3 which shows that the linear combination of predictors are equal to the *log-odds*, also called *logits*.

Once estimates of the model parameters β_0 and β_1 have been calculated, predictions of the class probability can be made using the logistic link function:

$$\hat{p}(X) = \sigma(\hat{\beta}_0 + \hat{\beta}_1 X) = \frac{e^{\hat{\beta}_0 + \hat{\beta}_1 X}}{1 + e^{\hat{\beta}_0 + \hat{\beta}_1 X}}$$

From there, a *threshold value* can be used to predict class membership, for example based on $\hat{p}(X) > 0.5$, but other decision rules and threshold values may be used, depending on the desired proportion or cost of false positives and true positives.

The binary logistic regression can be extended to *multinomial logistic regression*, where there are more than two classes for the output or target. The form of the equations is similar for K classes, starting from the log-odds. In this derivation, the last class K plays the role of reference class. As classes are not ordered, this choice is arbitrary.

$$\log\left(\frac{\Pr(Y = k|X = x)}{\Pr(Y = K|X = x)}\right) = \beta_{k0} + \beta_{k1}x_1 + \cdots + \beta_{kp}x_p, k < K$$

Exponentiating and multiplying:

$$\Pr(Y = k|X = x) = \Pr(Y = K|X = x)e^{\beta_{k0} + \beta_{k1}x_1 + \cdots + \beta_{kp}x_p}, k < K \quad (10.4)$$

Because probabilities must sum to 1:

$$\Pr(Y = K|X = x) = 1 - \sum_{l=1}^{K-1} \Pr(Y = l|X = x)$$

Substituting Eq. 10.4 into the right-hand side:

$$= 1 - \sum_{l=1}^{K-1} \Pr(Y = K|X = x)e^{\beta_{l0} + \beta_{l1}x_1 + \cdots + \beta_{lp}x_p}$$

Moving $\Pr(Y = K|X = x)$ out of the sum, dividing by it, and rearranging:

$$\Rightarrow \Pr(Y = K|X = x) = \frac{1}{1 + \sum_{l=1}^{K-1} e^{\beta_{l0} + \beta_{l1}x_1 + \cdots + \beta_{lp}x_p}} \quad (10.5)$$

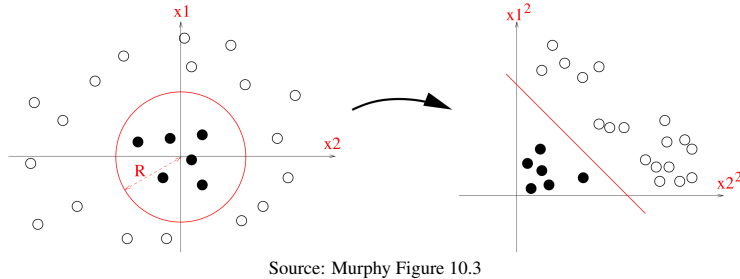


Figure 10.14: Transforming non-linear decision boundaries using polynomials

Substituting Eq. 10.5 into Eq. 10.4:

$$\Rightarrow \Pr(Y = k | X = x) = \frac{e^{\beta_{k_0} + \beta_{k_1}x_1 + \dots + \beta_{k_p}x_p}}{1 + \sum_{l=1}^{K-1} e^{\beta_{l_0} + \beta_{l_1}x_1 + \dots + \beta_{l_p}x_p}}, k < K \quad (10.6)$$

Equations 10.5 and 10.6 give the class probabilities for the reference class K and for any other class $k < K$. They are formally similar to the equations for the binary logistic regression above; for example, compare equation 10.6 to equation 10.1.

Just as polynomials of the input variables may be useful as predictors in linear regression, they also have applications in classification. In particular, they transform non-linear decision boundaries into linear ones that can be modeled using linear logistic regression. For example, the nonlinear boundary in the left panel of Figure 10.14 can be transformed into a linear boundary by squaring both predictors, as shown in the right panel of Figure 10.14.

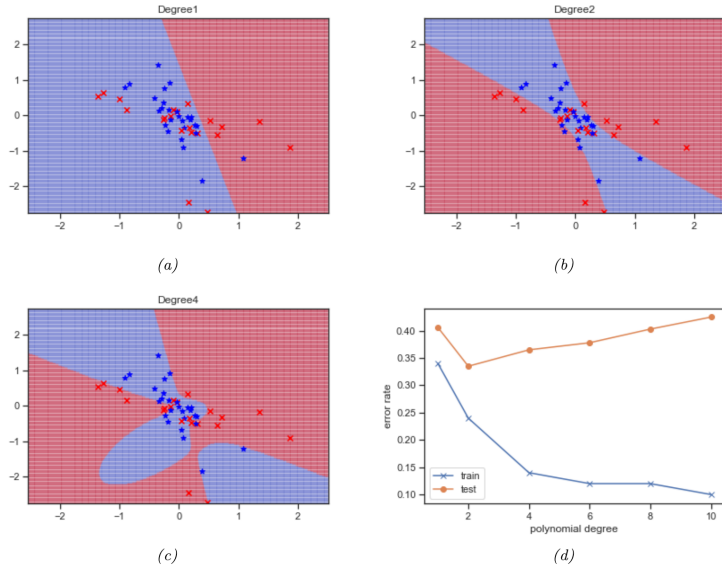
Another example is shown in Figure 10.15 where the linear decision boundary dividing the top left panel into a blue and red region clearly does not fit the observations, shown as blue and red points. Fitting logistic regressions with different degrees of polynomials shows that the decision boundary can be transformed to better fit the observed data. The bottom right panel in Figure 10.15 shows the the train and test error rate for different degrees of polynomials.

10.7.2 Logistic Regression in R

The following illustration of logistic regression in R uses the Smarket data set of the ISLR2 library. The data set contains stock market information and is used in this example to predict the direction of the movement of the market, either "up" or "down", based on previous day's ("lagged") data and other variables⁴.

Logistic regression can be performed using the same `glm()` function as for linear regression; it is one specific form of the generalized linear model. The `family` argument is used to indicate the type of regression and the link function:

⁴The R code for this example is based on material in Section 4.7.2 of ISLR2



Source: Murphy Figure 10.4

Figure 10.15: Transforming linear decision boundaries using polynomials

```
library(ISLR2)
?Smarket

# Contrasts show how factor levels are encoded using dummy variables:
contrasts(Smarket$Direction)

# Fit a logistic regression model
logreg.fitted <-
  glm(Direction~Lag1+Lag2+Lag3+Lag4+Lag5+Volume, data=Smarket,
       family=binomial(link='logit'))
summary(logreg.fitted)
```

The `predict(.)` function for the fitted model can be used to predict either the logits or the class probabilities:

```
# Predict logits for training data
logreg.logits <- predict(logreg.fitted, newdata = Smarket)

# Predict probabilities for training test
logreg.probabilities <- predict(logreg.fitted, newdata = Smarket,
                                 type='response')
```

A decision rule is necessary to assign observations to classes using the calculated proba-

bilities. The following example classifies them into the "Up" class, if its class-membership probability is greater than .5:

```
# Predict 'up' or 'down' based on probabilities and a threshold
pred.direction <- rep('Down', nrow(Smarket))
pred.direction[logreg.probabilities > .5] <- 'Up'
```

The confusion matrix can be produced by using the `table()` function on the predicted class and the observed class. Accuracy is simply the average number of observations for which predicted class and observed class are identical.

```
# Compute confusion matrix
logreg.cm <- table(pred.direction, Smarket$Direction)
print(logreg.cm)

# Compute accuracy
mean(pred.direction == Smarket$Direction)
```

The next R code blocks illustrate the use of a holdout sample or validation set approach to evaluate the performance of the logistic regression classifier.

Because the data set is a time series, a random split is not appropriate because it would mean that the training set would contain information later in time that informs the model parameter estimates which are used to predict earlier observations in the test set. Instead, time series data must be split non-randomly at some point in time:

```
train.data <- Smarket[Smarket$Year < 2005,]
test.data <- Smarket[!(Smarket$Year < 2005),]
```

Next, the model is fitted to the training data set, class-membership probabilities for observations in the test data set are then predicted, and the observations are classified using a decision rule:

```
logreg.fitted <-
  glm(Direction~Lag1+Lag2+Lag3+Lag4+Lag5+Volume, data=train.data,
       family=binomial(link='logit'))

# Predict probabilities for test data and classify:
logreg.probabilities <- predict(logreg.fitted, newdata = test.data,
                                 type='response')
pred.direction <- rep('Down', nrow(test.data))
pred.direction[logreg.probabilities > .5] <- 'Up'
```

While the confusion matrix is useful to understand how a classifier behaves, it is only a first step in evaluating the classifier performance. The ROCR library provides the `performance()` function to evaluate the predictive performance of a classifier. It

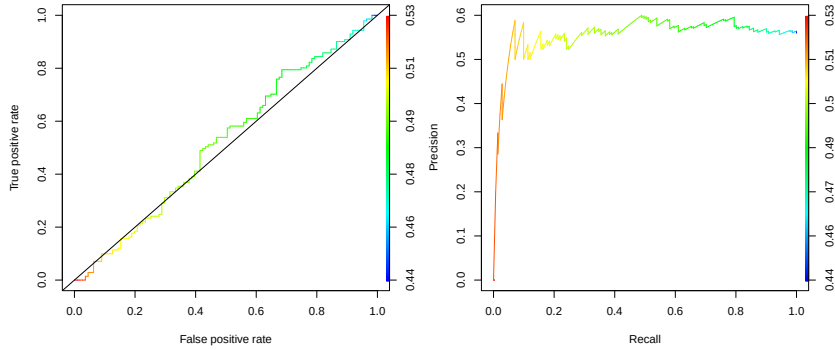


Figure 10.16: ROC and precision/recall curves for a logistic regression classifier

provides metrics such as accuracy, precision, recall, and can generate ROC curves. The last two plots generated by the R code block below are shown in Figure 10.16. The left panel shows the ROC curve, the right panel shows the precision/recall plot. It is clear from the results that predicting the stock market from the inputs in the given data set does not work well.

```
library(ROCR)

# A prediction object contains probabilities and true labels
pred.obj <- prediction(logreg.probabilities, test.data$Direction)

# Get some classifier performance metrics, ROCR varies the threshold.
plot(performance(pred.obj, 'acc'))
plot(performance(pred.obj, 'prec'))
plot(performance(pred.obj, 'rec'))
plot(performance(pred.obj, 'f'))

# ROC curve: True positive rate versus false positive rate
plot(performance(pred.obj, 'tpr', 'fpr'), colorize=T)
abline(0, 1)
# Precision/Recall plot
plot(performance(pred.obj, 'prec', 'rec'), colorize=T)
# Calculate the AUC
performance(pred.obj, 'auc')@y.values[[1]]
```

10.7.3 Naive Bayes Classifier

The naive Bayes classifier is based on *Bayes' theorem* of conditional probabilities. The probability that an observation described by a vector of inputs X is a member of class c can be described as the probability of observing inputs X given that the class is c , multiplied by the unconditional probability of an observation being in class c , divided

by the unconditional observation of observing inputs X . Formally:

$$\Pr(Y = c | X) = \frac{p(X|Y = c)p(Y = c)}{p(X)}$$

The overall probability of observing vector X is the sum of the probabilities of observing X in each class l , multiplied by the probability of an observation being member of class l :

$$= \frac{p(X|Y = c)p(Y = c)}{\sum_{l=1}^K p(X|Y = l)p(Y = l)} \quad (10.7)$$

The *naive Bayes assumption* is that within each class c , the D different input features that make up X are independently distributed of each other. With this assumption, one can write:

$$\begin{aligned} p(X|Y = c) &= p(x_1|Y = c) \times p(x_2|Y = c) \times \cdots \times p(x_D|Y = c) \\ &= \prod_{d=1}^D p(x_d|Y = c) \end{aligned} \quad (10.8)$$

Substituting Equation 10.8 into Equation 10.7 yields the *posterior probability* of class membership:

$$p(Y = c|X) = \frac{\left(\prod_{d=1}^D p(x_d|Y = c)\right) p(Y = c)}{\left(\sum_{l=1}^K \prod_{d=1}^D p(x_d|Y = l)\right) p(Y = l)}$$

The probabilities in the product of Equation 10.8 can be trivially estimated from the data, simply as the proportion of each x_d for each class c . However, the assumption of independence is violated when features are correlated. In other words, the naive Bayes classifier can be expected to perform best for independent features.

10.7.4 Naive Bayes Classifier in R

The e1071 library for R provides the `naiveBayes(.)` function that implements the naive Bayes classifier. The following illustration uses the same `Smarket` data from the `ISLR2` library that was used in the above example on logistic regression⁵:

⁵The R code for this example is based on material in Section 4.7.5 of ISLR2

```

library(e1071)
library(ISLR2)
train.data <- Smarket[Smarket$Year < 2005,]
test.data <- Smarket[!(Smarket$Year < 2005),]

# Fit using same syntax as glm
nb.fitted <- naiveBayes(Direction ~ Lag1 + Lag2, data=train.data)
# Output contains prior and conditional probabilities (and their SD)
print(nb.fitted)

```

A `predict(.)` method is available to predict class membership, given a fitted naive Bayes classifier and a data set of observations. A confusion matrix can be constructed by comparing predicted classes to observed classes. The following R code predicts class memberships and computes the confusion matrix for the test data.

```

nb.predictions <- predict(nb.fitted, test.data)
nb.cm <- table(nb.predictions, test.data$Direction)
print(nb.cm)

```

Evaluating the classifier is similar to evaluating the logistic regression classifier and again uses the ROCR library. The ROC curve created by the following R code block is shown in Figure 10.17. Comparing this ROC curve to the one in Figure 10.16 shows that the naive Bayes classifier performs slightly better than the logistic regression classifier.

```

library(ROCR)

# Predict probabilities (for use with ROCR)
nb.probabilities <- predict(nb.fitted, test.data, type='raw')
# Create an ROCR prediction object
nb.pred.obj <- prediction(nb.probabilities[, 'Up'], test.data$Direction)

# Generate an ROC plot
plot(performance(nb.pred.obj, 'tpr', 'fpr'), colorize=T)
abline(0, 1)
# Compute the AUC value
performance(nb.pred.obj, 'auc')@y.values[[1]]

```

10.7.5 KNN Classification

K-Nearest Neighbour classification is a non-parametric classification technique. It identifies the k nearest neighbours of a new observation, typically using the Euclidean distance or the L2-norm of the vector differences. It then estimates the class membership probabilities as the class membership proportions among the k nearest neighbours of the new observations. Details can be found in the previous chapter.

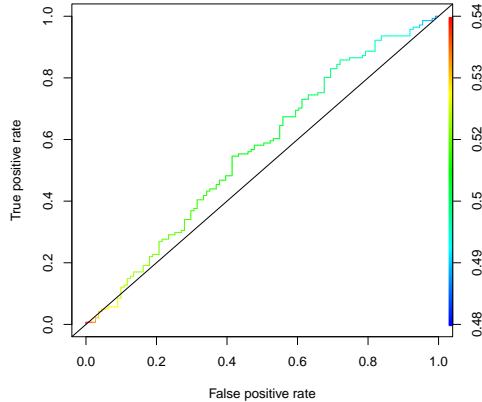


Figure 10.17: ROC curve of a naive Bayes classifier

One important requirement in KNN classification is *scaling of inputs* to have similar standard deviations or variance to avoid the distance metric being dominated by the input on the largest scale. For example, if one input ranges between 1 and 10 million, while another input ranges between 1 and 10, the first input is clearly most important in determining the distance between two observations.

10.7.6 KNN Classification in R

The `class` library for R provides the `knn(.)` function, as illustrated in the following example R code block that uses the same `Smarket` stock market data set as the above examples for logistic regression and naive Bayes⁶.

```
library(class)
library(ISLR2)
train.data <- Smarket[Smarket$Year < 2005,]
test.data <- Smarket[!(Smarket$Year < 2005),]

# Split the data into test and train sets
train.x <- cbind(train.data$Lag1, train.data$Lag2)
test.x <- cbind(test.data$Lag1, test.data$Lag2)
train.y <- train.data$Direction
test.y <- test.data$Direction
```

Next, the `knn(.)` function is used to make predictions for the test data set, given the training data inputs and training data outputs. The `knn(.)` function uses the Euclidean distance metric. The following R code example considers $k = 3$ near-

⁶The R code for this example is based on material in Section 4.7.6 of ISLR2

est neighbours and returns the class membership probabilities in addition to the class memberships of the test data set observations:

```
knn.pred <- knn(train.x, test.x, train.y, k=3, prob=T)
```

A confusion matrix can be computed using the `table(.)` function and the accuracy is calculated as the mean number (proportion) of observations for which the knn prediction is the same as the observed class membership.

```
# Confusion matrix
table(knn.pred, test.y)
# Accuracy
mean(knn.pred == test.y)
```

The class membership probabilities returned in the `knn(.)` function result are those of the majority class, in this example the class "Down". Because this is a binary classification, the class membership probabilities for the "Up" class can be trivially calculated, as shown in the following R code block:

```
knn.probs <- attributes(knn.pred)$prob

# Compute class probabilities of the minority class:
knn.class.probs <- knn.probs
knn.class.probs[knn.pred=='Down'] <- 1-knn.probs[knn.pred=='Down']
```

With the class membership probabilities for both classes, the ROCR library functions can be used to evaluate the classifier by plotting the ROC curve and computing the AUC. The ROC curve produced by the R code block below is shown in Figure 10.18.

```
knn.pred.obj <- prediction(knn.class.probs, test.data$Direction)
plot(performance(knn.pred.obj, 'tpr', 'fpr'), colorize=T)
abline(0, 1)
performance(knn.pred.obj, 'auc')@y.values[[1]]
```

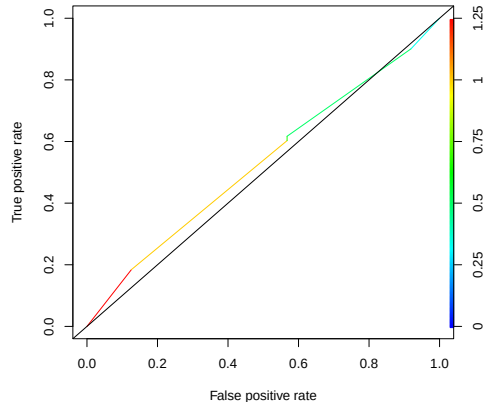


Figure 10.18: ROC curve of a k-NN classifier

Hands-On Exercise

Use the `Weekly` data set in the `ISLR2` package.

1. Use the full data set to perform a logistic regression with `Direction` as target. Which predictors are statistically significant?
2. Compute the confusion matrix and accuracy.
3. Use the 1990 to 2008 data for a training set and the 2009/2010 for a test set. Fit a logistic regression model with `Lag2` as the only predictor.
4. Repeat (3) using Naive Bayes
5. Repeat (3) using KNN with $K = 1$
6. Which model provides the best results on this data?

Source: ISLR2 Section 4.8

Hands-On Exercise

Use the `Auto` data set in the `ISLR2` package.

1. Create a binary variable, `mpg01` that contains a 1 if `mpg` is above its median, 0 otherwise. *Tip:* Use the `median()` function. Add the new variable to the data frame.
2. Split the data set into training and test set
3. Perform a logistic regression on the training data to predict `mpg01` from the other features. What is the test error of this model?
4. Repeat (3) using Naive Bayes
5. Repeat (3) using KNN with different values of K . What value of K performs best?

Source: ISLR2 Section 4.8

Hands-On Exercise

Using the `Boston` data set in the `ISLR2` library, fit classification models to predict whether a given census tract has a crime rate above or below the median.

1. Create a new binary variable `crime01` that is 1 if `crime` is above its median, and 0 otherwise. Combine this variable with the data frame. *Tip:* Use the `median()` function for this.
2. Split your data set into a training and test data set
3. Fit logistic regression, Naive Bayes, and KNN (with different K)
4. Describe your findings in terms of prediction error, precision, recall, F1 and AUC

Source: ISLR2 Section 4.8

10.8 Review Questions

Linear Regression

1. Explain the primary objective of linear regression and how it is implemented in a statistical model.
2. Discuss the importance of visual inspection of data before choosing a regression model.
3. Define *fitted values* or *predicted values* in the context of linear regression. How are they computed?
4. Discuss why a model with a term like $\beta_2 X^2$ is still considered a linear regression model.
5. How does the residual sum of squares (RSS) relate to mean squared error (MSE) in linear regression analysis?
6. Explain the importance and use of the *standard errors* of the estimates in linear

regression.

7. Describe what a *t-test* in regression analysis involves, and how it is used to test hypotheses about model parameters.
8. What does the R^2 statistic tell us about a linear regression model? What are its limitations?
9. Explain the term *interaction effects* using an example, and describe how they can be identified in a regression model.
10. What role do dummy variables play when incorporating categorical predictors into a regression model? Give an example.
11. How can the inclusion of more predictors into a linear regression model affect the model's bias and variance?

Random Numbers

12. Define a pseudo-random number generator (RNG). How does it differ from a true random number generator?
13. Discuss the role of the seed in the generation of pseudo-random numbers. What happens if the seed is not set before generating random numbers in a program?
14. Describe a scenario where using the same seed value might be advantageous in computational analyses.

Shrinkage Methods

15. Explain what is meant by "shrinkage methods" in the context of regression analysis. Why is it necessary to shrink the magnitude of regression coefficients?
16. What is the difference between L1 and L2 regularization? Provide examples where each might be preferable.
17. Explain the rationale behind using ridge regression and LASSO as alternatives to standard linear regression. What problem do they address?
18. Discuss why it is important to standardize predictors before applying ridge regression. What could happen if the predictors are on different scales?
19. Explain the concept of the LASSO as a form of penalized regression. How does it differ from ridge regression in terms of the impact on model parameters?
20. Discuss the method of selecting the penalty parameter λ in shrinkage methods like ridge regression and LASSO.
21. Discuss how the bias-variance trade-off is managed in ridge regression through the adjustment of λ . What are the signs that λ is set too high or too low?
22. Explain how the Elastic Net method balances the properties of L1 and L2 penalties. What role does the parameter α play in this balance?

Logistic regression

23. Explain the concept of the sigmoid or logistic function as a solution for bounding the output of a regression model between 0 and 1.
24. What is a link function in logistic regression? Describe its purpose and how it modifies the output of a linear model.

25. Define the logistic function and explain how it is used in logistic regression to estimate probabilities.
26. In logistic regression, what does the logit (or log-odds) function represent? How does it relate to the probabilities of class memberships?
27. Discuss the significance of the threshold value in logistic regression. How is it used to determine class membership?
28. Explain the concept of multinomial logistic regression and how it extends the binary logistic regression model to multiple classes.
29. Discuss how incorporating polynomial terms of input variables into a logistic regression model can help in transforming non-linear decision boundaries into linear ones.

Naive Bayes classification

30. What is Bayes' theorem and how is it applied in the naive Bayes classifier to compute class probabilities?
31. Explain how the probability $p(X|Y = c)$ is calculated under the naive Bayes assumption.
32. Discuss the implications of the independence assumption among features in the naive Bayes classifier. What are the potential limitations of this assumption in real-world scenarios?
33. Explain the role of prior probabilities $p(Y = c)$ in the naive Bayes classifier. How do these influence the final classification?
34. What is the effect of having highly correlated features on the performance of the naive Bayes classifier?

KNN classification

35. Define K-Nearest Neighbor (KNN) classification. Why is it classified as a non-parametric technique?
36. Explain the concept of "nearest neighbors" in the context of KNN. What metrics can be used to determine proximity in feature space?
37. Discuss the role of the number k in KNN classification. How does the choice of k influence the classifier's performance?
38. Describe how KNN estimates the class membership probabilities for a new observation.
39. Discuss the impact of feature scaling on the performance of KNN classification. Why is it important to scale features?
40. Discuss the trade-offs between choosing a larger versus smaller value of k in KNN classification.

Chapter 11

Introduction to Unsupervised Machine Learning

Sources and Further Reading

The material in this chapter is based on the following sources. They are freely available. Consult them for additional information.

Gareth James, Daniel Witten, Trevor Hastie and Robert Tibshirani: *An Introduction to Statistical Learning with Applications in R*. 2nd edition, corrected printing, June 2023. (ISLR2)

<https://www.statlearning.com>

Chapter 12

The book by James et al. provides an easy introduction to machine learning at the introductory undergraduate level. It focuses on applications, not mathematics, and contains many exercises using R. Concepts are well explained and illustrated. There is a similar book available by the same authors with applications in Python. This book is a more accessible of the following book.

Trevor Hastie, Robert Tibshirani, and Jerome Friedman: *The Elements of Statistical Learning*. 2nd edition, 12th corrected printing, 2017. (ESL)

<https://hastie.su.domains/ElemStatLearn/>

Chapter 14

The book by Hastie et al. still sets the standard for statistical learning. It is widely used and cited. Its treatment is more technical than the previous book and there are

no exercises in R or Python. However, it covers the concepts in more depth (and a few more formulas). However, it is still very accessible even to an undergraduate audience.

Kevin P. Murphy: *Probabilistic Machine Learning – An Introduction*. MIT Press 2022.

<https://probml.github.io/pml-book/book1.html>

Chapters 20, 21

Murphy's book is available under a creative-commons license. It is a somewhat more technical treatment of the material, but with many illustrations and examples. It is quite comprehensive in its coverage and targeted at the advanced undergraduate or graduate student.

11.1 Introduction

In unsupervised machine learning, there are no known correct outputs that can be used to train or fit statistical models. In that sense, there are no X and Y variables, but only the X variables. Unsupervised machine learning focuses on identifying patterns in the data, often in order to simplify the data. The two unsupervised methods considered here, principal components analysis (PCA) and cluster analysis or clustering, both do this. For example, PCA "summarizes" multiple variables or "dimensions" into fewer variables, the principal components, while cluster analysis finds similarities in the data and groups or clusters observations into fewer clusters than observations. The principal components and the clusters can be viewed as simplifications or summaries of the data.

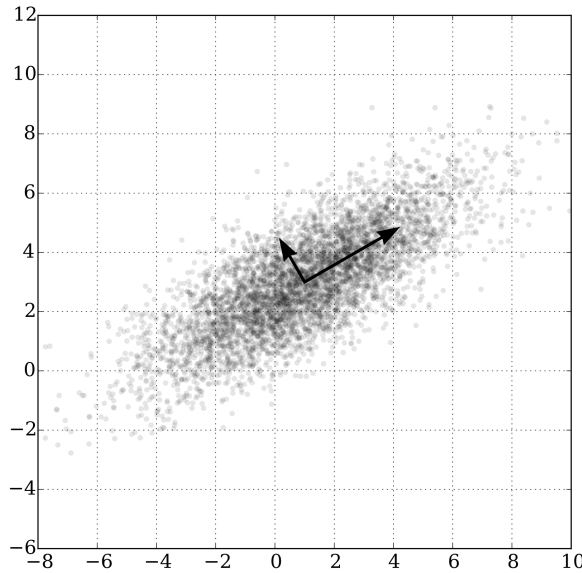
11.2 Principal Components Analysis

The aim of *principal component analysis* (PCA) is to create linear combinations of the input variables, the principal components (PC), that satisfy two conditions:

1. They are maximally variable, that is, their variance is maximal, and
2. They are orthogonal, that is, independent, of each other.

There are as many principal components as there are input variables. Generally, only a few of the principal components, the ones with the greatest variance, are retained for further analysis. It is not uncommon to reduce hundreds of variables to five or ten principal components for further analysis.

These principal components are considered summaries of the original data and can be used, for example, instead of the original input variables in a regression or classification model. This makes the model smaller and therefore easier to understand, interpret, and verify. Using fewer inputs for a regression or classification can also serve as a regularization method, that is, a way to make the model less susceptible to overfitting. This is because models with fewer inputs generally have fewer parameters, all other



<https://commons.wikimedia.org/wiki/File:GaussianScatterPCA.svg>

Figure 11.1: Scatterplot with Principal Components

things being equal. Additionally, the principal components are useful for data visualization. It is much easier to show a 2D or 3D summary of the data when the data has been summarized in two or three principal components, rather than visually depicting dozens or hundreds of variables.

Figure 11.1 shows an example visualization of a scatterplot of data on two variables and the two principal components. Technically, the two arrows shown are the *eigen-vectors* of the covariance matrix of the data, scaled in length by the square root of the corresponding *eigenvalue* and then shifted to the mean of the data, details that will become clear below.

We first introduce an iterative method of computing principal components. Recall that principal components are linear combinations of the original input variables. Hence, the first principal component (PC) for $1 \leq i \leq n$ data values and p variables is defined as:

$$z_{i1} = w_{11}x_{i1} + w_{21}x_{i2} + \cdots + w_{p1}x_{ip}$$

Or, simpler, in matrix form:

$$Z_1 = Xw_1$$

The *weight vector* or *loading vector* $w_1 = (w_{11}, \dots, w_{p1})$ is a $p \times 1$ column vector that is scaled to unit length, that is, $\|w_1\|_2 = 1$. X is a $n \times p$ data matrix and Z_1 is the first principal component of size $n \times 1$.

Assuming zero-centered variables, the variance of Z_1 and the optimization criterion can be expressed as follows:

$$\underset{w_{j1}}{\text{maximize}} \sum_{i=1}^n z_{i1}^2 = \sum_{i=1}^n \left(\sum_{j=1}^p w_{j1} x_{ij} \right)^2 \quad (\text{Variance of } z_{i1}) \quad (11.1)$$

Subject to:

$$\sum_{j=1}^p w_{j1}^2 = 1 \quad (\text{Scaling constraint})$$

Or, simpler, in matrix form:

$$\underset{w_1}{\text{maximize}} Z_1^T Z_1 = w_1^T X^T X w_1 \quad (\text{Variance of } Z_1) \quad (11.2)$$

Subject to:

$$\|w_1\|_2 = 1 \quad (\text{Scaling constraint})$$

To derive the second PC, subtract the first PC from the data:

$$X_{\text{new}} \leftarrow X - X w_1 w_1^T$$

Then, repeat the maximization with the residual data X_{new} , that is the "left over" portion of the data.

This procedure can be repeated until as many principal components k are calculated as there are original data variables p . Because each iteration reduces the remaining data, the residual, by subtracting a component with maximum variance, the variance of the residual data shrinks. Hence, the variance of each successive principal component and therefore the proportion of the initial overall variance accounted for by each successive principal component shrinks. In other words, each successive component explains a decreasing proportion of the total original variance in the data.

There are two important considerations when working with PCA. First, the input data variables should be scaled to have equal or unit standard deviation, so that the measurement scale of different variables does not influence the outcome of the PCA. Second,

the signs of the principal components can be "flipped" arbitrarily. This can be seen in Figure 11.1, where one can easily imagine the two arrows pointing in opposite direction, and still providing the same good summary of the original data.

Variables in the data set should be scaled to identical standard deviations prior to PCA.

To give an applied example, consider four input variables extracted from a data set of police arrest data in the US for violent crimes in each of the 50 states of the US. While four principal components can be computed, the four input variables can be summarized pretty well by just the first two principal components that together explain more than 80% of the total variance. Figure 11.2 shows a plot of the data along the first two components which form the horizontal and vertical axis. This is known as a *biplot*. Overlaid are the four original variables. Table 11.1 shows the *component loadings*, that is the ϕ in the above formulas, for the first two principal components.

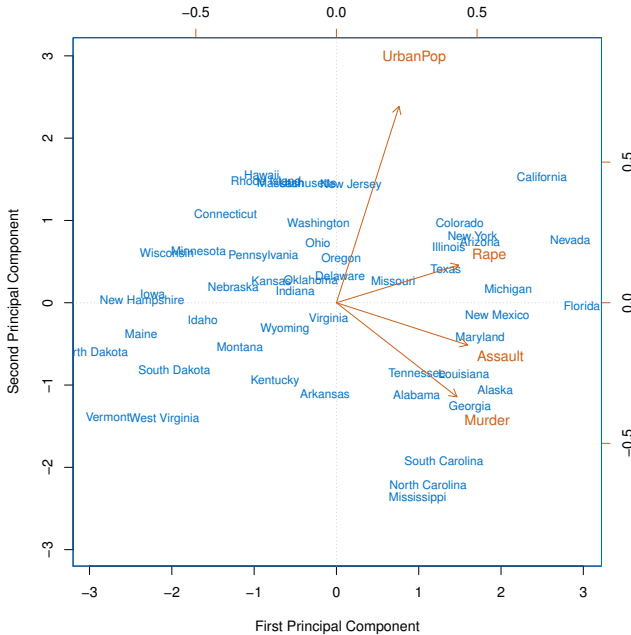
Interpretation of the principal components, which is important in explanation but less so in prediction, focuses on the loadings. For example, looking at the columns of the loadings in Table 11.1 shows that the first PC has high loadings on the variables "Murder", "Assault", and "Rape", and a much smaller loading on "UrbanPop". This suggests that PC1 expresses the overall prevalence of violent crime, as a summary of those three variables. In contrast, the second PC has a high loading on "UrbanPop", but a much lower (absolute) loading on the other three variables, indicating that it expresses primarily the one variable "UrbanPop". This interpretation is supported by Figure 11.2, which examines the rows of Table 11.1, plotting each variable as a two-component vector in the space spanned by PC1 and PC2 (recall that the principal components are by definition orthogonal). Here, the row vector for the variable "UrbanPop" is visually distinct and separate from the row vectors for the other three variables.

While the iterative description of principal components above illustrates the properties of the components in terms of their variance, actual PCA is done by means of *eigen-decomposition*. It turns out that the solution to the maximization problem in Equations 11.1 and 11.2 are the principal components of the data correlation matrix. Each principal component is an *eigenvector* of the data correlation matrix such that:

	PC1	PC2
Murder	.536	-0.418
Assault	.583	-0.188
UrbanPop	.278	0.873
Rape	.543	0.167

Source: ISLR2 Table 12.1

Table 11.1: US arrest data example – first two principal component loadings



Source: ISLR2 Figure 12.1

Figure 11.2: US arrests data example – Biplot with data plotted on first two principal components with original variables

$$V^{-1}X^T XV = V^{-1}CV = \Lambda$$

where V is the matrix whose columns are the *eigenvectors*, C is the data correlation matrix, and Λ is a diagonal matrix of *eigenvalues*.

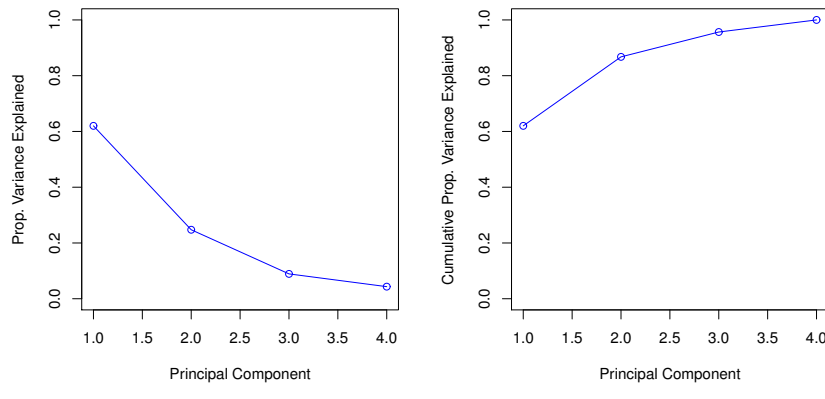
The *proportion of variance explained* f_k by each PC k is proportional to the corresponding *eigenvalue* λ_k , that is, the k -th entry of Λ :

$$f_k = \frac{\lambda_k}{\sum_{j=1}^p \lambda_j}$$

The *cumulative proportion of variance* F_k explained by the first k PC is then:

$$F_k = \frac{\sum_{j=1}^k \lambda_j}{\sum_{j=1}^p \lambda_j}$$

There are different criteria for selecting the number of principal components to retain for further analyses:



Source: ISLR2 Figure 12.3

Figure 11.3: US arrests data example – Scree plot and cumulative variance explained

- There may be a theoretical reason, especially in an explanation context, to retain a specific number of principal components
- The analyst retains those principal components that have an intuitive and relevant interpretation, as in the above example.
- The analyst retains those principal components whose eigenvalue $\lambda > 1$
- The analyst retains principal components until the cumulative proportion of variance explained by the components surpasses a given threshold, e.g. 80%. For example, the right panel in Figure 11.3 shows a plot of the cumulative variance explained. The first two principal components are necessary to explain 80% or more of the total variance in the original data.
- When used in subsequent regression or classification models, cross-validation may be used to identify the optimal K that shows the lowest test error.
- The analyst examines the "scree plot", that is, the plot of the eigenvalues or proportion of variance explained by each component. Oftentimes, there will be a clear point of inflection in this plot, indicating a useful cutoff. The left panel in Figure 11.3 shows such a scree plot. The proportion of variance explained diminishes for each additional principal component.

In practice, the number of principal components to retain is often subjective, and analysts use a combination of considerations and criteria to make their decision.

11.3 Principal Components Analysis in R

The `USArrests` in the `ISLR2` library contains data on the arrests (per 100,000 residents) for various violent crimes as well as the percentage of urban population in the

50 states of the US¹. First, examine the data and the correlation between variables. In the correlation matrix, one can already see that "UrbanPop" is not highly correlated with the other three variables, an indication that it will not load on the same principal component as those.

```
library(ISLR2)
?USArrests
summary(USArrests)
cor(USArrests)
```

The `prcomp()` function in R performs a PCA and can optionally scale and center the data before doing so:

```
# PCA using prcomp()
# Scaling is generally a good idea
pca.result <- prcomp(USArrests, scale=TRUE)

# Print the component loadings
pca.result$rotation
```

The results can be plotted in a biplot, similar to the one in Figure 11.2, using the `biplot()` function for the `prcomp` result object. By default, that function uses the first two principal components, but others can be specified using the `choices` argument. Note that the signs of the principal components may be arbitrarily flipped.

```
# Biplot for components 1 and 2
biplot(pca.result, choices=1:2, scale=0)
```

The explained variance can be computed from the result and plotted in a scree plot similar to the one in the left panel of Figure 11.3.

```
# Explained variance for each component
pca.result$sdev^2

# Scree plot (both points and lines)
plot(pca.result$sdev^2, type='b', col='blue')
```

Recall that the proportion of variance explained is the proportion of the variance of a principal component out of the total variance explained by all principal components. The R function `cumsum()` can be used to conveniently calculate the cumulative value of this. The following code block computes a cumulative plot similar to the right panel in Figure 11.3.

¹The R code for this example is based on material in Section 12.5 of ISLR2

```
# Proportion of variance explained
pve <- pca.result$sdev^2 / sum(pca.result$sdev^2)

# Cumulative sum of variance explained
plot(cumsum(pve), type='b', col='blue')
```

Using the `eigen(.)` function for eigenvalue decomposition shows that the principal component loadings correspond to the eigenvectors and the explained variance corresponds to the eigenvalues,

```
# Eigen-decomposition of correlation matrix
e <- eigen(cor(USArrests))
# Compare values and vectors to prcomp results
e$values
e$vectors
```

The component scores themselves are also available in the `prcomp` result for use in further analysis such as regression or classification:

```
# Print the component scores themselves
# For further use in regression, etc.
head(pca.result$x)
```

Hands-On Exercise

The Boston dataset in the `ISLR2` library describes house prices in the different suburbs of Boston. Use PCA to reduce the number of dimensions for this dataset:

1. Use the `prcomp` function to perform a PCA on the centered and standardized data. Limit yourself to quantitative inputs.
2. Produce a biplot of the first two components
3. Provide the proportion of variance explained by each component
4. How many components would you retain? Why? How much of the total variance would this explain?
5. Based on the loadings, can you ascribe meaning to the components? What do they represent?

Hands-On Exercise

The `Harmann74.cor` dataset in the `datasets` library contains the results of 24 psychological tests given to 145 school children. Use PCA to reduce the number of dimensions for this dataset:

1. Use the `prcomp` function to perform a PCA on the centered and standardized data. Limit yourself to quantitative inputs.
2. Produce a biplot of the first two components
3. Provide the proportion of variance explained by each component
4. How many components would you retain? Why? How much of the total variance would this explain?
5. Based on the loadings, can you ascribe meaning to the components? What do they represent?

Hands-On Exercise

The `Hitters` dataset in the `ISLR2` library contains the salary of 322 baseball players and season statistics. Use `salary` as the target variable and all other numerical variables as predictors.

1. Use PCA to reduce the number of dimensions for the predictors. Limit yourself to quantitative inputs.
2. Retain the first principal component.
3. Estimate and cross-validate a regression model using the first PC as predictor. What is the training and validation error?
4. Repeat steps (1) to (3), retaining 2, 3, ..., all components
5. Plot the training and validation error against the number of components. Describe and discuss your results.

11.4 Clustering

Whereas PCA tried to simplify a data set "by column" through the identification of variables that can be summarized by principal components, cluster analysis tries to simplify a data set "by row" through the identification of observations that are similar and can be represented as a group, that is, a *cluster*. The aim is to form homogenous subgroups of observations and to discover "structure" in the data.

There are many different types of clustering. This chapter focuses on two simple and easy-to-understand methods. The *k-means clustering algorithm* is an example of centroid-based clustering, a method that assigns observations to clusters based on their distance from the cluster center ("centroid"), while *agglomerative clustering* is a form of hierarchical clustering which iteratively merges observations together to form larger and larger clusters.

11.4.1 K-Means Clustering

In k-means clustering, the number of clusters K is assumed given, determined by the analysts knowledge of the data or the requirements of the analysis. The aim of k-means clustering is to minimize the *within-cluster variation* $W(C_i)$ in each cluster C_i :

$$\min_{C_i} \left\{ \sum_{k=1}^K W(C_k) \right\}$$

This within-cluster variation is defined as the squared Euclidean distance between every pair of observations in the cluster (Equation 11.3) or between every observation and the cluster *centroid* of the cluster it is assigned to, that is, its corresponding cluster mean $\bar{\mu}$ (Equation 11.4).

$$W(C_k) = \frac{1}{|C_k|} \sum_{i,i' \in C_k} \sum_{j=1}^p (x_{ij} - x_{i'j})^2 \quad (11.3)$$

$$= 2 \sum_{i \in C_k} \sum_{j=1}^p (x_{ij} - \bar{\mu}_{kj})^2 \quad (11.4)$$

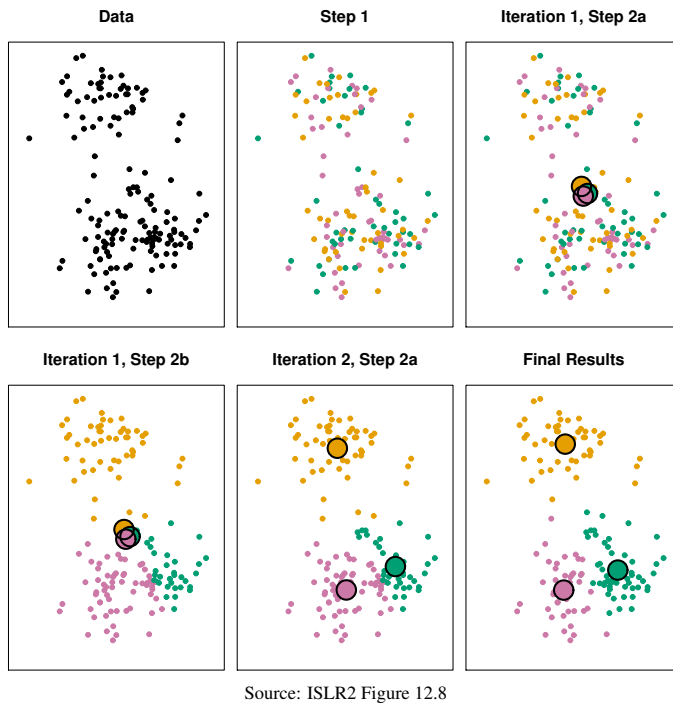
Here, i, i' range over observations within cluster C_k , j ranges over the p different variables that make up an observation, and $\bar{\mu}_{kj}$ is the mean of variable j for cluster k .

This definition of distance means that k-means cluster analysis is only applicable to quantitative variables.

When variables are measured on different scales, e.g. one variables in the range of $[0, 1]$ while another is measured between $[0, 1000000]$ it is important to *standardize or scale the variables to have similar standard deviations* (typically, unit standard deviation, i.e. 1). Otherwise, the Euclidean distance between observations is dominated by the variable with the largest range.

Variables in the data set should be scaled to identical standard deviations prior to k-means clustering.

K-means clustering uses an iterative algorithm, beginning with the random assignment of each observation i to one of the k clusters. From these cluster assignments, the cluster means (centroids) can be computed ($\bar{\mu}_{kj}$ in Equation 11.4). Next, each observation is assigned to that cluster whose centroid is closest. The last two steps are repeated until the cluster assignments no longer change.

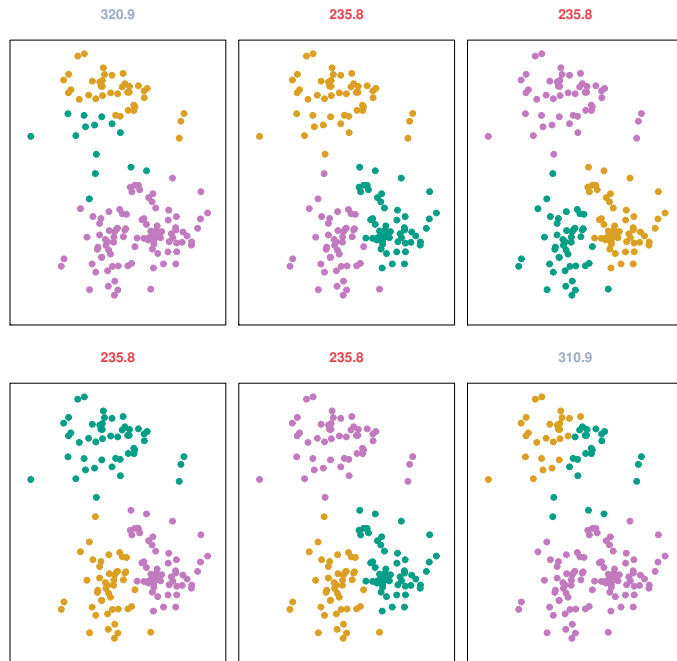


Source: ISLR2 Figure 12.8

Figure 11.4: K-means iterative cluster assignment example

This process is illustrated in Figure 11.4. The top left panel shows observations on two variables. The panel labeled "Step 1" shows the initial random assignment of each observation to one of three clusters, indicated by the color. The top right panel, labelled "Iteration 1, Step 2a" shows the cluster means or centroids computed based on this assignment as large coloured circles. As one might imagine, random assignment leads to cluster means that are very similar. The bottom left panel, "Iteration 1, Step 2b" shows the cluster assignment of the observations based on the new cluster means. Each observation is assigned to that cluster whose mean is closest. The bottom middle panel, "Iteration 2, Step 2a" shows the new cluster means based on the new cluster assignment of observations. The bottom right panel shows the final, stable cluster assignment. Repeated calculation of cluster means and assigning observations to clusters does not change cluster membership for any observation. Note that the cluster membership in this final panel is slightly different than the one in the bottom middle panel, indicating at least one more iteration between the two panels.

It should be clear from this description that the random initial cluster assignment has a significant impact on the final result. As the number of observations grow, the random effects generally diminish, but different random initial cluster assignments may yield different final clustering solutions.



Source: ISLR2 Figure 12.9

Figure 11.5: K-means clustering solutions from different initial cluster assignments

The k-means algorithm should be run multiple times and the optimal solution, that is, the one with the lowest *within-cluster variability*, should be chosen for further analysis.

This effect is shown in Figure 11.5. The data from the previous example was clustered six different times with different random initial cluster assignments. Each final solution is different, and may also have a different *within-cluster variability* as shown at the top of each panel. Note that some solutions are identical but permute the cluster assignments/colours. For example, the top middle and top right panels in Figure 11.5 are identical and also identical with the bottom left and bottom middle solution, except for the permutation of cluster assignments, indicated by the colours.

11.4.2 K-Means Clustering in R

To illustrate k-means clustering in R, consider the following simulated example, which uses the `kmeans()` function². Data is simulated as 50 observations on two normally distributed variables. One half of the data is shifted by +3 on the first variables and

²The R code for this example is based on material in Section 12.5 of ISLR2

350CHAPTER 11. INTRODUCTION TO UNSUPERVISED MACHINE LEARNING

by -4 on the second variable. With a standard deviation of 1, this constitutes a large separation and should lead to clearly identifiable clusters.

```
# Set RNG seed for replicability
set.seed(2)
# Create a 50 x 2 matrix of random variables
# Normally distributed, with 0 mean and SD=1
x <- matrix(rnorm(n=50*2, mean=0, sd=1), ncol=2)
# Clearly separate the first 25 points by shifting their coordinates
x[1:25, 1] <- x[1:25, 1] + 3
x[1:25, 2] <- x[1:25, 2] - 4
```

Next, the data is clustered using the `kmeans()` function into 2 clusters, 20 times with different random initial cluster assignments:

```
# Cluster into 2 clusters, performing 20 random starting assignments
km.result <- kmeans(x, 2, nstart=20)
```

The result object `km.result` contains the cluster means, the cluster assignments for each observation and the sum-of-squares (distances) within each cluster and between clusters. Recall that the optimization objective is to minimize the within-cluster variation.

```
# Results show cluster means, cluster assignments,
# and sums of squares (distances) within and between
print(km.result)
# Those values are also available as components in the result object
names(km.result)
print(km.result$centers)
print(km.result$withinss)
# etc.
```

Finally, it is easy to create colour-coded plot of the data (the following R code block adds 1 to every cluster number to avoid plotting black points). This generates a plot as shown in Figure 11.6, clearly indicating the well-separated clusters.

```
# Plot the color-coded points
plot(x, col=(km.result$cluster+1),
      main = 'K-Means Clustering Results with K=2',
      xlab = '', ylab='', pch=20, cex=2)
```

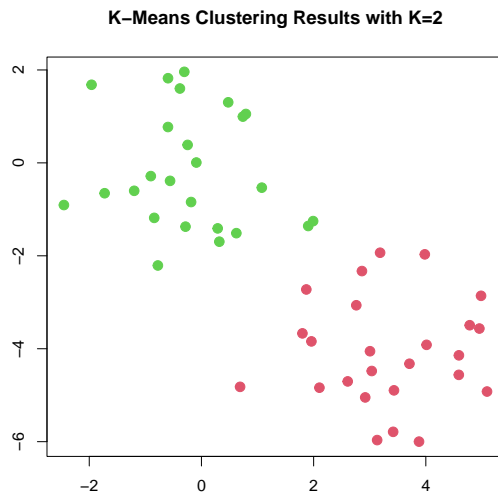


Figure 11.6: Result of k-means clustering on simulated data

Hands-On Exercise

The Boston dataset in the `ISLR2` library describes house prices in the different suburbs of Boston. Use K-Means Clustering to identify sets of similar suburbs using only the numerical variables in the data set.

1. Use the `kmeans` function to perform a cluster analysis, using multiple starting assignments. Limit yourself to quantitative inputs but do not scale the variables.
2. Use different numbers of clusters k and identify which value of k gives you the best results. Define what you mean by "best" and justify your choice.
3. Scale the data so that each variable has the same variance or standard deviation, but do not change the variable means.
4. Repeat the cluster analysis with the best value of k and compare results.

Hands-On Exercise

The `Hitters` dataset in the `ISLR2` library contains the salary of 322 baseball players and season statistics. Use K-Means Clustering to identify sets of similar players, using only the numerical variables in the data set.

1. Use the `kmeans` function to perform a cluster analysis, using multiple starting assignments. Limit yourself to quantitative inputs but do not scale the variable.
2. Use different numbers of clusters k and identify which value of k gives you the best results. Define what you mean by "best" and justify your choice.
3. Scale the data so that each variable has the same variance or standard deviation, but do not change the variable means.
4. Repeat the cluster analysis with the best value of k and compare results.

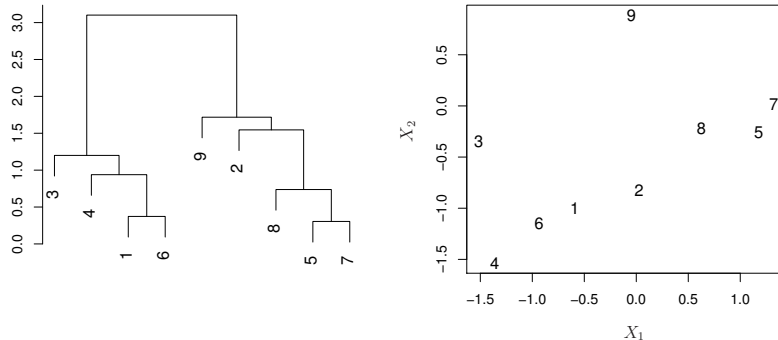
11.4.3 Hierarchical Clustering

Hierarchical clustering is either *agglomerative*, that is, it constructs clusters "bottom-up" by joining observations or small clusters to larger clusters, or it may be *divisive*, that is, in "top-down" fashion, starting from the whole set of observations, it iteratively divides the set into clusters. This section examines the use of agglomerative clustering, which is widely used because of its intuitive process and its flexibility.

Agglomerative clustering begins with n observations and a distance (or, alternatively, a similarity metric, which is just the inverse of distance – a large distance means a small similarity). The process is then as follows:

1. Treat each observation as its own cluster
2. Repeat the following steps $n - 2$ times:
 - (a) Calculate distances between all pairs of clusters
 - (b) Identify the pair of clusters that are least distant from each other
 - (c) "Fuse" or merge these two clusters

The process is usually visualized with a *dendrogram*, which literally means "tree graph", such as the one shown in the left panel of in Figure 11.7. A dendrogram is read bottom-up, showing which clusters are merged in which order. The vertical axis shows the distance between clusters as they are merged. Consider the observations on two variables shown in the right panel of Figure 11.7. In the example, clusters 5 and 7 are merged first, from a distance of ≈ 0.3 . This distance is the smallest distance between all clusters, indicated as the lowest merging point in the dendrogram in the left panel of Figure 11.7. Cluster 5 is just observation 5, and cluster 7 is just observation 7. The two together form a new cluster. Next, clusters 1 and 6 are merged, from a distance of ≈ 0.4 , the second lowest merging point in the dendrogram. Then, cluster 8 (which is observation 8) is added to the cluster consisting of observations 5 and 7, at a distance



Source: ISLR2 Figure 12.12

Figure 11.7: Example dendrogram and data for agglomerative clustering

of ≈ 0.8 . After this, observation 4 is added to the cluster consisting of observations 1 and 6, etc. The final two clusters are at a distance of ≈ 3 when they are merged into a single cluster.

The following key decisions need to be made by the analyst for agglomerative clustering:

- How to measure similarity or distance between observations?
- How to measure distance between clusters ("linkage")?
- How many clusters should there be?

Table 11.2 shows a set of common distance metrics or vector norms that are frequently used in agglomerative clustering. Figure 11.8 is a visualization of the intuition behind some of these distance metrics. For example, the Chebyshev distance allows diagonal "moves" to count as a single step with a distance of 1, whereas the taxicab metric counts a "move" in each direction as a single step, so that diagonal "moves" have a distance of 2. In principle, any of these distance metrics could also be used in k-means clustering, but this is rarely done.

Because the distance function is heavily influenced by the measurement scale of the variables, when these are not equal, it is possible for one variable to dominate others, simply because it is measured on a different scale. As with PCA and k-means clustering, it is therefore important to scale the variables in the data set to identical standard deviation (typically, unit standard deviation, i.e. 1).

Variables in the data set should be scaled to identical standard deviations prior to hierarchical clustering.

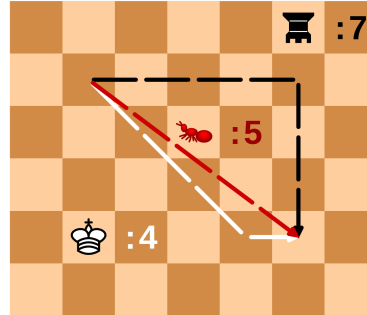
Table 11.3 shows a set of the most commonly used *linkage functions*, that is, functions that express the distance between two clusters G and H . The *single linkage* is based on the minimum distance of any pair of observations where one observation is in cluster

Taxicab or Manhattan	$\ q - p\ _1$	$\sum_i q_i - p_i $
Euclidean	$\ q - p\ _2$	$\sqrt{\sum_i (q_i - p_i)^2}$
Minkowski	$\ q - p\ _p$	$\left(\sum_i q_i - p_i ^p\right)^{\frac{1}{p}}$
Chebyshev	$\ q - p\ _\infty$	$\lim_{p \rightarrow \infty} \left(\sum_i q_i - p_i ^p\right)^{\frac{1}{p}} = \max_i(q_i - p_i)$
	$\ q - p\ _{-\infty}$	$\lim_{p \rightarrow -\infty} \left(\sum_i q_i - p_i ^p\right)^{\frac{1}{p}} = \min_i(q_i - p_i)$

Table 11.2: Common distance metrics or "norms" in clustering

$1 \ 1 \ 1$	$\sqrt{2} \ 1 \ \sqrt{2}$	$2 \ 1 \ 2$
$1 \ \text{♚} \ 1$	$1 \ \text{LOBSTER} \ 1$	$1 \ \text{TOWER} \ 1$
$1 \ 1 \ 1$	$\sqrt{2} \ 1 \ \sqrt{2}$	$2 \ 1 \ 2$

Chebyshev Euclidean Taxicab



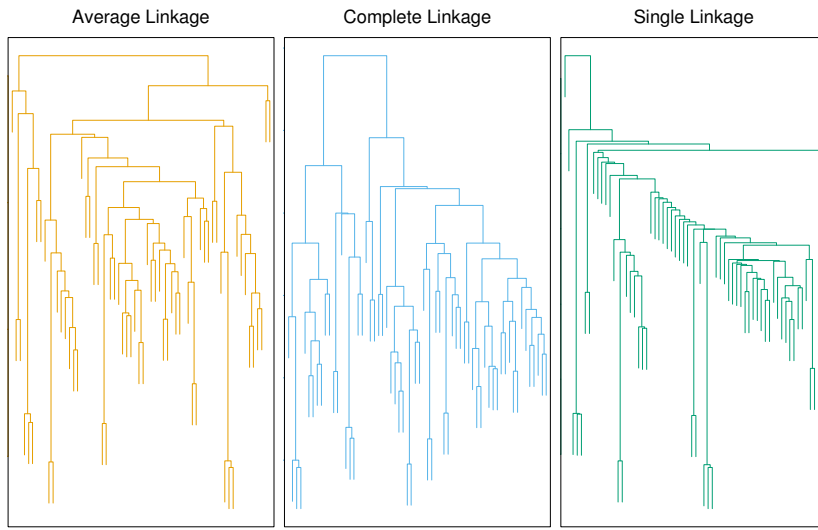
https://commons.wikimedia.org/wiki/File:Minkowski_distance_examples.svg

Figure 11.8: Different distance metrics and their intuition

G and the other in cluster H . In other words, the distance of two clusters is the distance between the two closest observations from each cluster. In contrast, *complete linkage* uses the maximum; the distance between clusters is the maximal distance between any of their member observations. Finally, *average linkage* uses the mean distance between all pairs of observations. There are many other, less commonly used linkage functions

Single	$d_{SL}(G, H) = \min_{i \in G, i' \in H} d_{i, i'}$
Complete	$d_{CL}(G, H) = \max_{i \in G, i' \in H} d_{i, i'}$
Average	$d_{AL}(G, H) = \text{mean}_{i \in G, i' \in H} d_{i, i'}$

Table 11.3: Commonly used linkage functions in hierarchical clustering



Source: ISLR2 Figure 12.14

Figure 11.9: The effect of different linkage functions in agglomerative clustering

available³.

The linkage function has a significant effect on the process of clustering a set of observations. Consider the three examples shown in the different panels of Figure 11.9. Merging two observations into a cluster is always done at the same distance, as this is determined purely by the distance metric, not the linkage function. However, the decision which clusters (of multiple observations) to combine is heavily influenced by the linkage function as can be seen in the very different dendograms in Figure 11.9.

The final question concerns the choice of the number of clusters. The answer to this question may be driven by theory (typically in explanatory applications), by requirements of the subsequent data analysis or the subsequent use of the resulting clusters, or by examining the distances at which clusters are merged, that is, the height in the dendrogram. Choosing a number of clusters is called "cutting the dendrogram" at a specific point. Consider the example in Figure 11.10. The left panel shows the solution of the agglomerative clustering. In the end, a single cluster containing all the

³https://en.wikipedia.org/wiki/Hierarchical_clustering

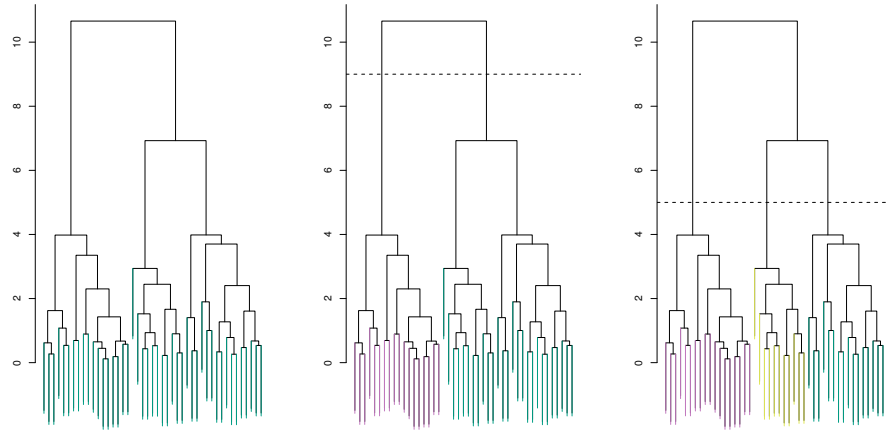


Figure 11.10: Cutting a dendrogram to determine the number of clusters

observations remains, with the last two clusters merged at a distance of ≈ 10.5 . The middle and right panel show two different "cuts" of the dendrogram, one resulting in two clusters and the other resulting in three clusters. The cuts may be determined by a desired number of clusters, by considerations of distance, or both. It should be clear that lowering the "cut" height further beyond what is shown in the right panel, that is reducing the distance between clusters, would result in many small clusters with a much smaller distance between them.

11.4.4 Hierarchical Clustering in R

This example uses the same simulated data as the example for k-means clustering⁴. First, generate 50 observations on two variables from a normal distribution. One half of the observations are shifted on both variables to provide a known cluster structure.

```
# Set RNG seed for replicability
set.seed(2)
# Create a 50 x 2 matrix of random variables
# Normally distributed, with 0 mean and SD=1
x <- matrix(rnorm(n=50*2, mean=0, sd=1), ncol=2)
# Clearly separate the first 25 points by shifting their coordinates
x[1:25, 1] <- x[1:25, 1] + 3
x[1:25, 2] <- x[1:25, 2] - 4
```

The `dist()` function is used to calculate differences between the observations. The names for the `method` argument to `dist()` are the same as in Table 11.2. Additionally, the '`maximum`' distance in R uses the greatest distance among all the variables

⁴The R code for this example is based on material in Section 12.5 of ISLR2

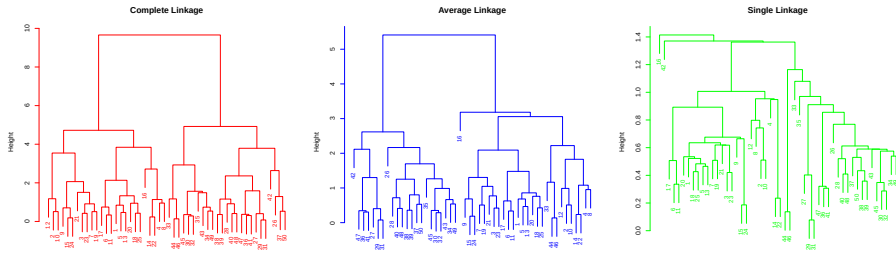


Figure 11.11: Dendrogram of three clustering solutions for simulated data

of the two observations.

```
# The dist() function calculated distances
# according to a variety of metrics/norms
euclid.dist <- dist(x, method='euclidean')
pnorm.dist <- dist(x, method='minkowski', p=3)
manh.dist <- dist(x, method='manhattan')
max.dist <- dist(x, method='maximum')
```

The `hclust()` function performs the hierarchical agglomerative clustering. The `method` argument specifies the type of linkage, according to Table 11.3. The `hclust()` function can use a few additional linkages not listed in that table, see the documentation (`?hclust`) for details.

```
# Use the hclust() function with a distance metric
hc.complete <- hclust(euclid.dist, method='complete')
hc.single <- hclust(euclid.dist, method='single')
hc.average <- hclust(euclid.dist, method='average')
```

The dendograms for the three different clustering solutions can be plotted to produce Figure 11.11.

```
# Plot the dendograms in a single plot
par(mfrow = c(1, 3))
plot(hc.complete, col='red',
      main = "Complete Linkage", xlab = "", sub = "", cex = .9)
plot(hc.average, col='blue',
      main = "Average Linkage", xlab = "", sub = "", cex = .9)
plot(hc.single, col='green',
      main = "Single Linkage", xlab = "", sub = "", cex = .9)
```

The complete linkage and average linkage solutions are visually quite similar, but upon careful examination of which observations and clusters are merged in which order,

they are actually very different from each other. The single linkage solution is visually very different from the others. Note the "height" of the dendrogram on the vertical axis. Because the single linkage focuses on the minimal distance between a pair of observations from each cluster, the heights in this dendrogram are the smallest among the three dendograms. Because the average linkage focuses on the mean of distances of pairs of observations of two clusters, its height values are generally larger, but still smaller than the range of heights for the complete linkage solution, which focuses on the maximum distance between pairs of observations from two clusters.

Finally, cutting the dendrogram is done with the `cutree()` function by specifying either the number of clusters k or the height h at which the dendrogram is to be cut. The function returns a vector with the cluster membership for each observation.

```
# Cut by number of groups/clusters
cutree(hc.complete, k=4)
# Cut by height (distance)
cutree(hc.complete, h=6)
```

Hands-On Exercise

The Boston dataset in the `ISLR2` library describes house prices in the different suburbs of Boston. Use Hierarchical Clustering to identify sets of similar suburbs using only the numerical variables in the data set.

1. Use the `hclust` function to perform a cluster analysis, exploring different distance metrics and linkage functions. Limit yourself to quantitative inputs.
2. Examine the dendograms and identify which combination of distance metric and linkage function gives you the "best" solution. Define "best" and justify your decision.
3. How many clusters k would you choose?
4. Using this value for k , perform a k-means Clustering and compare the results. Remember that k-means clustering uses the Euclidean distance.

Hands-On Exercise

The `Hitters` dataset in the `ISLR2` library contains the salary of 322 baseball players and season statistics. Use Hierarchical Clustering to identify sets of similar players, using only the numerical variables in the data set.

1. Use the `hclust` function to perform a cluster analysis, exploring different distance metrics and linkage functions. Limit yourself to quantitative inputs and make sure you scale the data.
2. Examine the dendograms and identify which combination of distance metric and linkage function gives you the "best" solution. Define "best" and justify your decision.
3. How many clusters k would you choose?
4. Using this value for k , perform a k-means clustering and compare the results. Remember that k-means clustering uses the Euclidean distance.

Hands-On Exercise

The `Auto` dataset in the `ISLR2` library contains information on 392 vehicles. Use Hierarchical Clustering to identify sets of similar vehicles, using only the numerical variables in the data set.

1. Use the `hclust` function to perform a cluster analysis, exploring different distance metrics and linkage functions. Limit yourself to quantitative inputs.
2. Examine the dendograms and identify which combination of distance metric and linkage function gives you the "best" solution. Define "best" and justify your decision.
3. How many clusters k would you choose?
4. Using this value for k , perform a k-means Clustering and compare the results. Remember that k-means clustering uses the Euclidean distance.

11.5 Review Questions

Principal Components Analysis

1. Explain how unsupervised machine learning differs from supervised machine learning in terms of data requirements and outcomes.
2. What are the main goals of Principal Component Analysis (PCA) in data analysis?
3. Explain the concept of "variance" in the context of PCA. Why is maximizing variance an important objective?
4. How can PCA be used to simplify a complex dataset? Give an example based on a hypothetical dataset.
5. How can PCA contribute to improving the interpretability of complex models?

6. Describe the process of calculating the first principal component in PCA. What role do the loading vectors play? What optimization problem does PCA solve?
7. How does one interpret the loadings of a principal component and what do they signify about the variables involved?
8. Discuss the importance of scaling input variables before performing PCA. What could potentially happen if the variables are not scaled?
9. Describe the relationship between eigenvalues and the variance explained by the principal components. How does one interpret these eigenvalues in practical terms?
10. Provide several criteria that could be used to decide how many principal components to retain in an analysis.
11. Discuss the relevance of the "scree plot" in determining the number of principal components to retain. What does an inflection point in the scree plot typically indicate?
12. Explain how the biplot can be used to visualize both the principal components and the original variables. What insights can one gain from such a visualization?
13. Explain how PCA can be used as a feature extraction technique in machine learning models.

K-Means Clustering

14. Define clustering in the context of unsupervised machine learning and explain its main purpose.
15. Compare and contrast the goals of principal component analysis (PCA) and clustering.
16. What are centroid-based clustering and hierarchical clustering? Provide examples of each.
17. Describe the k-means clustering algorithm. What objective does it aim to achieve?
18. Explain the concept of within-cluster variation in the context of k-means clustering.
19. What are the implications of variable scales on the performance of the k-means clustering algorithm? Why might scaling be necessary?
20. Illustrate the iterative process of the k-means clustering algorithm. What happens in each step?
21. Explain why the initial random assignment of observations to clusters can affect the final clustering solution in k-means.
22. Discuss the importance of running the k-means algorithm multiple times. How does this practice influence the reliability of the clustering results?
23. What are the computational complexities of k-means and hierarchical clustering? How do these affect their scalability to large datasets?
24. Discuss the limitations of k-means clustering and possible scenarios where it might not perform well.

Hierarchical Clustering

25. Describe a scenario in which hierarchical clustering would be more beneficial

- than k-means clustering. Consider aspects such as data structure and analysis goals.
26. Describe hierarchical clustering and differentiate between agglomerative and divisive clustering.
 27. Explain the initial steps in an agglomerative clustering process. How does it begin, and what happens in the initial stages?
 28. Define a dendrogram and explain how it is used in hierarchical clustering.
 29. Discuss the significance of distance measures in hierarchical clustering. How do they affect the clustering process?
 30. How might the concept of distance be adapted when clustering categorical data using hierarchical methods?
 31. What are the different types of linkage methods in hierarchical clustering? Describe at least three and explain how they influence the clustering results.
 32. Provide an overview of common distance metrics used in agglomerative clustering. How might the choice of distance metric influence the outcome of clustering?
 33. Explain the process of creating a dendrogram and interpreting its structure in the context of hierarchical clustering.
 34. Explore the relationship between the number of observations and the interpretability of the dendrogram in hierarchical clustering. How does increasing the number of observations affect the clarity and usefulness of the dendrogram?
 35. Explain the concept of "cutting the tree" in hierarchical clustering. How does this process determine the number of clusters?
 36. Discuss how the choice of linkage method might impact the sensitivity of hierarchical clustering to outliers and noise in the dataset.
 37. How does the analyst decide on the number of clusters in hierarchical clustering? What factors might influence this decision?
 38. Consider the distance metrics shown in Table 11.2. Which metric would be most appropriate for clustering data with extreme outliers and why?
 39. Explain why it might be necessary to standardize variables before performing hierarchical clustering.
 40. Evaluate the computational complexity of hierarchical clustering. How does this complexity influence the scalability of the method to large datasets?

Chapter 12

Time Series Analysis

Sources and Further Reading

The material in this chapter is based on the following sources. Consult them for additional information and details.

Robert H. Shumway and David S. Stoffer (2017) *Time Series Analysis and Its Applications*, 4th Edition. Springer.

<https://www.stat.pitt.edu/stoffer/tsa4/>

The book by Shumway and Stoffer provides a very comprehensive but also somewhat technical introduction to the subject of time series analysis. The authors have also published the `astsa` library for R to accompany their book. This library provides a number of data sets and functions for time series analysis.

Rob J. Hyndman and George Athanasopoulos (2018) *Forecasting: Principles and Practice*, 2nd edition. OTexts.

<https://otexts.com/fpp2/>

The book by Hyndman and Athanasopoulos is somewhat less technical in nature than the book by Shumway and Stoffer and also provides R code. The coverage of the two books also differs somewhat, but it is more accessible than Shumway and Stoffer for undergraduate students.

In addition to these books, there are a number of very useful tutorials available on the internet that can augment or summarize the material in the books. They are less focused

on theory and more focused on actually performing time series analysis.

- <https://github.com/nickpoison/tsa4>
- <https://a-little-book-of-r-for-time-series.readthedocs.io/en/latest/src/timeseries.html>
- <https://rc2e.com/timeseriesanalysis>
- <https://atsa-es.github.io/atsa-labs/chap-tslab.html>

12.1 Introduction

This section provides an introduction to time series analysis. Time series analysis is a complex topic with a multitude of different methods and techniques and this section can provide only a glimpse at the basic ideas and concepts.

Time series analysis is a set of statistical techniques that involve analyzing time-ordered data points or observations to extract meaningful statistics and other characteristics. This type of analysis is important across various fields such as economics, where it may be used to model unemployment rates, finance, where it may be used to model stock prices, social science, where it may be used to model high school graduation rates, natural sciences, where it may be used to model weather and climate trends, ecology, where it could be used to model animal population numbers, or epidemiology where it may be used to model the spread of epidemics. Understanding trends, cycles, and patterns over time can lead to useful insights and informed decision-making. Figure 12.1 shows an example of a basic time series of the quarterly earnings per share of a company.

At its core, a time series is a sequence of data points recorded at successive time intervals. The data is typically collected at uniform intervals – be it hourly, daily, monthly, or yearly. Time series analysis helps in understanding the inherent structure and functions that generate the series. It aims to model the underlying context of the data,

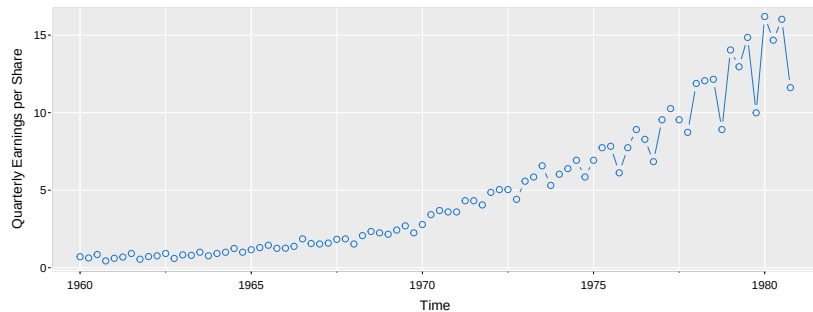


Figure 12.1: Example of time series data

whether to understand the past behavior or to forecast future values.

The analysis of time series can be divided broadly into two types: descriptive and inferential. Descriptive analysis focuses on visualizing and summarizing the main features of the data, such as trends (long-term direction), seasonality (regular pattern of fluctuation within a year), and irregular components (unpredictable, random fluctuations). Inferential analysis, on the other hand, involves using models to predict future values based on known past values, testing hypotheses, and deriving estimates of population parameters.

In time series analysis, two fundamental approaches to examining data are the time-domain and the frequency-domain approaches. The *time-domain approach* analyzes data as it evolves over time, focusing on the relationship between current and past values to predict future values. This approach is primarily concerned with understanding and modeling the temporal sequence directly in the time dimension. This approach is particularly useful for forecasting, where understanding how values are correlated through time is essential. It provides direct and often simple models that are interpretable in terms of the original time series data.

The *frequency-domain approach*, on the other hand, analyzes data based on the rate at which the data's features repeat over time. This approach transforms the time series data into the frequency domain using mathematical transformations (the most common being the Fourier Transform). It decomposes the time series into a combination of sinusoid functions with different frequencies and amplitudes. The frequency-domain approach is useful for identifying hidden periodicities or cyclical behaviors in the data, which may not be apparent in the time domain.

12.2 Time Series Statistical Models

Time series statistical models are essential tools used to analyze and forecast time-dependent data. Four common models are the moving average (MA) model, the autoregressive (AR) model, the random walk with drift, and the signal in noise model. Each model has different characteristics and applications, suited to different types of time series data.

Moving Average Model

The *Moving Average* model is a fundamental time series model that expresses the current value of the time series as a function of past errors or deviations, with the assumption that these errors are white noise, that is, random. An example model is given by:

$$v_t = \frac{1}{3}(w_{t-1} + w_t + w_{t+1})$$

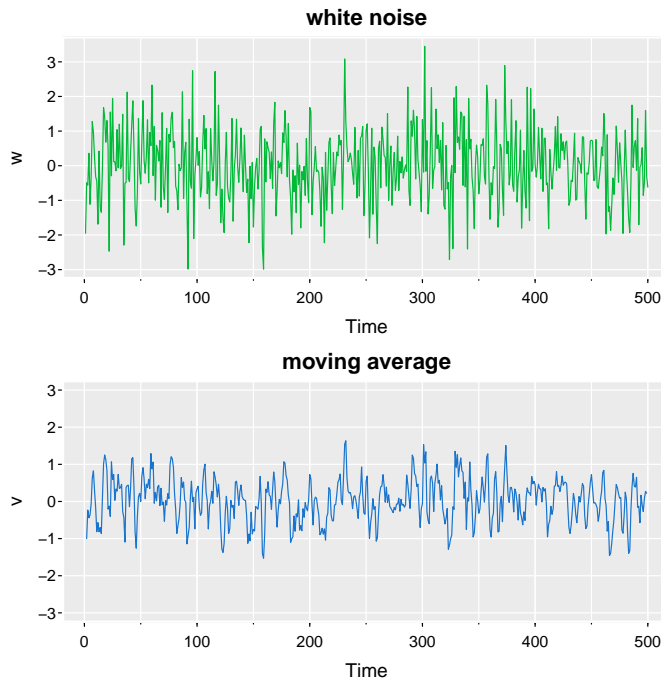


Figure 12.2: Example white noise time series and its moving average

where w_t are the white noise error terms. In this example, all white noise terms are weighted equally by $1/3$.

MA models are particularly useful in smoothing out noise and forecasting when the series exhibits a random behavior with no trend or seasonality.

The following R code block uses the `filter` function to generate the example model. The filter operates on the white noise, extending to both sides of the current time step, and creates a weighted sum of the closest three value in `w`, specified by the `mode='convolution'` argument. The resulting plots are shown in Figure 12.2.

```
# Random numbers as errors
w <- rnorm(500,0,1)
# Moving average
v <- filter(w, sides=2, filter=c(1/3,1/3,1/3), method='convolution')
# Plot timeseries
par(mfrow=c(2,1))
# The astsa library contains the tsplot function
library(astsa)
tsplot(w, main="white noise", col=3, gg=T)
tsplot(v, ylim=c(-3,3), main="moving average", col=4, gg=T)
```

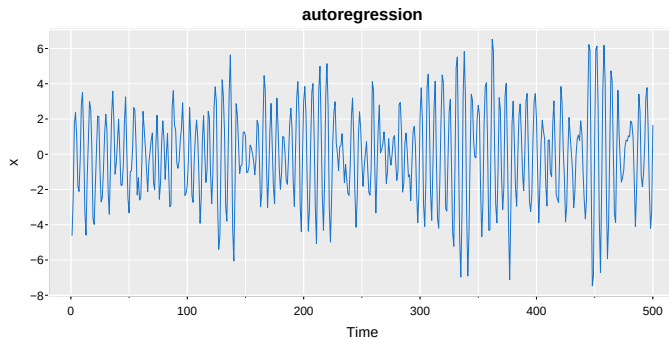


Figure 12.3: Example autoregressive time series

Autoregressive Model

The *Autoregressive (AR) model* is based on the concept that current values of a series can be forecasted from previous values. An example model is:

$$x_t = x_{t-1} - 0.9x_{t-2} + w_t$$

where w_t is white noise.

AR models are widely used in economic and financial time series where data points are influenced significantly by their previous values.

The following R code uses the `filter` function in "recursive" mode with parameters 1 and `- . 9` to create the time series corresponding to the example model¹. The resulting plot is shown in Figure 12.3.

```
# Random numbers (errors)
w <- rnorm(550,0,1)
# remove first 50 values for startup
x <- filter(w, filter=c(1,-.9), method="recursive") [-(1:50)]
tsplot(x, main="autoregression", col=4, gg=T)
```

Random Walk with Drift

A *random walk with drift* adds a constant to the standard random walk, allowing the series to drift upwards or downwards over time. An example model is given by:

¹The R code for this and following examples are based on material Shumway & Stoffer

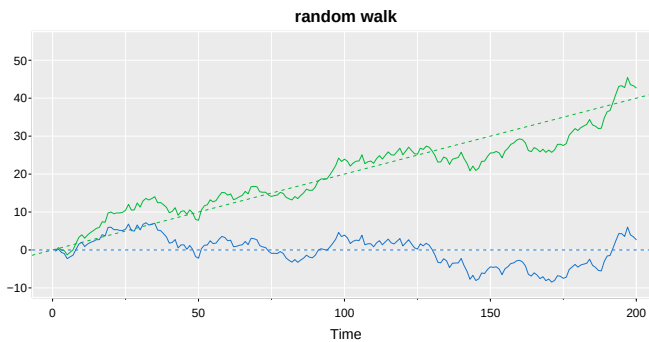


Figure 12.4: Example random walk with drift time series

$$\begin{aligned}x_t &= \delta + x_{t-1} + w_t \\&= \delta t + \sum_{j=1}^t w_j\end{aligned}$$

where δ represents the drift (constant term), and w_t is the noise component.

This model is commonly applied in financial markets to model stock prices or other investments, reflecting that prices are serially correlated and can trend over time.

The following R code block uses the the `cumsum()` function to calculate the cumulative sum. The resulting time series are shown in Figure 12.4 and show the random walk and the drift component that is added to it.

```
# Create random walk (white noise) model and then add drift
w <- rnorm(200)
x <- cumsum(w)
drift <- .2
w.drift <- w + drift;
x.drift <- cumsum(w.drift)
# Plot the two resulting series
tsplot(x.drift, ylim=c(-10,55), main="random walk", ylab='', col=3, gg=T)
abline(a=0, b=drift, lty=2, col=3)
lines(x, col=4)
abline(h=0, col=4, lty=2)
```

Signal in Noise Model

The *Signal in noise* model views the time series as a combination of a true signal and random noise. An example model with a sinusoidal signal characterized by its amplitude, frequency and phase shift is:

$$x_t = A \cos(2\pi\omega t + \phi)$$

for example,

$A = 2$	amplitude
$\omega = 1/50$	frequency
$\phi = .6\pi$	phase shift

This model is fundamental in signal processing and is used to understand underlying trends in the presence of noisy observations. Techniques like filtering and smoothing are often applied to extract the signal from x_t .

The following R code block creates a sinusoidal signal and overlays it with different amounts of white (Gaussian) noise. The resulting time series are shown in Figure 12.5.

```
# Create signal
cs = 2*cos(2*pi*1:500/50 + .6*pi)
w = rnorm(500,0,1)
# Overlay with gaussian noise and plot
par(mfrow=c(3,1), mar=c(3,2,2,1), cex.main=1.5)
tsplot(cs, main='Signal', col=2, gg=T)
tsplot(cs+w, main='Signal and N(0,1) noise', col=3, gg=T)
tsplot(cs+5*w, main='Signal and N(0,25) noise', col=4, gg=T)
```

12.3 Basic Time Series Operations in R

A time series can be constructed from an ordinary data set using the `ts()` function in R and supplying a start time stamp and a sampling frequency. For example, the following R code creates a time series of montly observations, beginning in January 2020 with the values 1 through 24. R will try to sensibly interpret the `start` and `frequency` arguments: Frequencies of 4 are interpreted as quarters of the year, 7 is interpreted as days of a week, 12 is interpreted as months of the year.

```
# Creating a time series object with monthly data
ts_data <- ts(1:24, frequency = 12, start = c(2020, 1))
```

To see the first and last last observations of a time series, use the `head()` and `tail()` functions:

```
head(ts_data)
tail(ts_data)
```

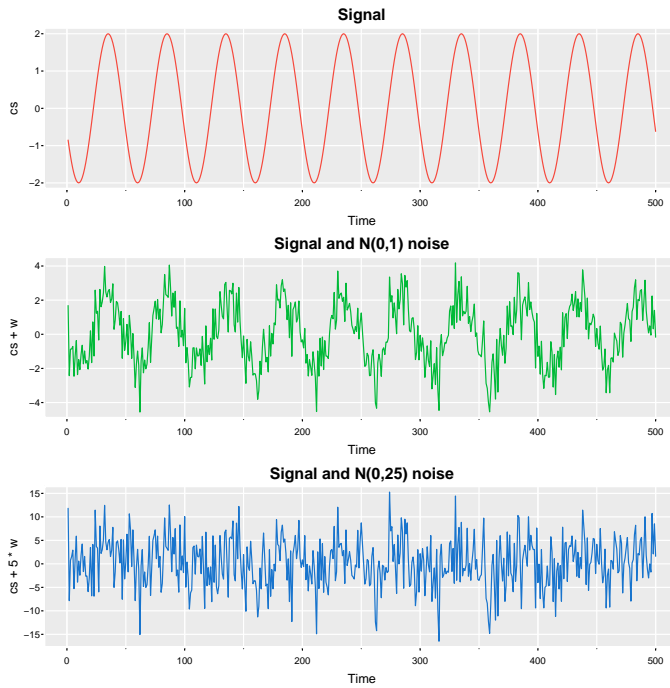


Figure 12.5: Example signal in noise time series

Missing values in a time series cannot be imputed in the usual manner because the observations are not independent of each other. Two simple ways of "filling in" missing data are either to simply carry the last observation forward, which assumes there are negligible changes in the value over time, or to interpolate the missing values. In linear interpolation, a line is imagined between the last observation before a series of missing values and the first observation after such a series of missing values. Missing values are then assumed to be on that line. This assumes that the time series is approximately linear, at least for short gaps.

The `zoo` library for R contains the functions `na.locf()` and `na.approx()` that implement these methods of handling missing values. Missing values at the beginning or end of a time series can be removed with the `na.trim()` function. The following R code block illustrates the use of all three functions:

```
# Introduce NA values into the time series
ts_data[c(5, 10, 15)] <- NA

# Using na.trim to remove leading/trailing NA values
trimmed_ts <- na.trim(ts_data)
# Using na.locf (Last Observation Carried Forward) to handle NA values
locf_ts <- na.locf(ts_data)
# Using na.approx to interpolate NA values
approx_ts <- na.approx(ts_data)
```

Two time series can be combined using the `ts.intersect()` or `ts.union()` functions. The former function combines the series only for overlapping, that is, intersecting, times, possibly cutting off the head or tail of one or the other series. The latter function retains all dates of both series and "pads" the head or tail of one or the other series with "NA" values.

```
# Creating another time series
ts_data2 <- ts(c(1:24), frequency = 12, start = c(2020, 7))

# Using ts.intersect to determine intersection of two time series
intersect_ts <- ts.intersect(ts_data, ts_data2)
# Using ts.union to determine union of two time series
union_ts <- ts.union(ts_data, ts_data2)
```

An important operation in time series analysis is to "lag" a time series, that is, to shift it forwards or backwards in time. R provides the `lag()` function for this purpose. A positive argument shifts the time series *backwards* by the specified number of time periods, while a negative argument shifts it forward:

```
# Positive k shifts backwards in time
lag_ts <- lag(ts_data, 2)
# Negative k shifts forwards in time
lag_ts <- lag(ts_data, 3)
```

12.4 Smoothing a Time Series

Time series smoothing is a technique used to remove noise and reveal signals or underlying trends in the data. Four commonly used methods for time series smoothing are moving average, kernel smoothing, lowess regression, and smoothing splines.

Moving Average Smoothing

Moving average smoothing is one of the simplest and most widely used methods for smoothing time series data. It involves calculating the weighted mean of the consecutive data points within a specified window that moves along with the data:

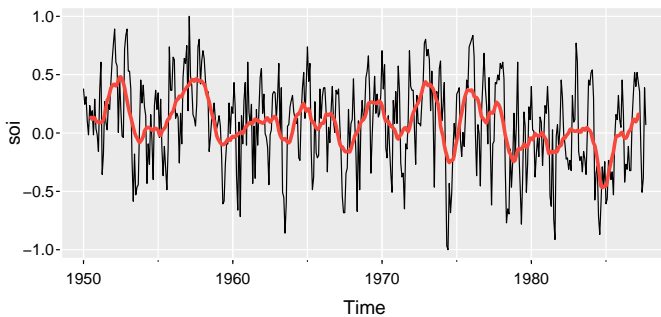


Figure 12.6: Moving average smoothing

$$m_t = \sum_{j=-k}^k a_j x_{t-j} \quad \text{where} \quad \sum_{j=-k}^k a_j = 1$$

where a are the weights that sum to one. A simple filter uses uniform weights, but other shapes are possible.

This model can be implemented using the `filter` function in R. The filter in the example below is two-sided and is centered on the current time stamp, that is, it uses data before and after the current time point. When the `sides=1` argument is used, the filter is over past values only. The `filter` argument specifies the weights for the moving average. The results for an example time series are shown in Figure 12.6.

```
# Use the soi dataset from the astsa library as an example
library(astsa)
?soi
# Apply moving average filter
f = 1/12 * c(0.5, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 0.5)
filter(soi, sides=2, filter=f, method='convolution')
```

Kernel Smoothing or Kernel Regression

Instead of a filter with simple weights as in moving average smoothing, *kernel smoothing* uses a weighted average of neighbouring points where the weights are determined by a function known as the *kernel*. A common choice is a Gaussian kernel that uses the normal distribution density function, which produces a weighted average with a bell-shaped curve of weights around each data point. The weights for averaging the time series values are determined as follows:

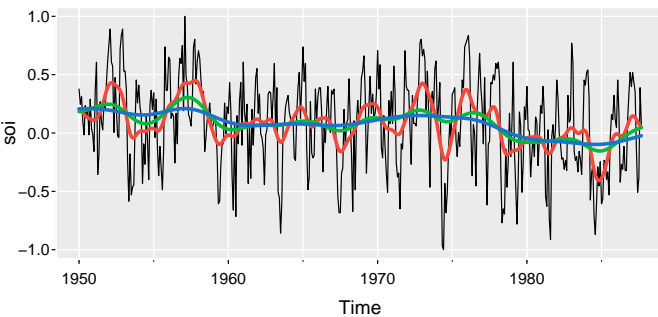


Figure 12.7: Kernel density smoothing with different kernel bandwidths

$$a_i(t) = \frac{K\left(\frac{t-i}{b}\right)}{\sum_{j=1}^n K\left(\frac{t-j}{b}\right)}$$

where K is the Gaussian kernel:

$$K(z) = \frac{1}{\sqrt{2\pi}} \exp(-z^2/2)$$

Here, b is the "bandwidth" of the kernel, that determines the shape of the kernel function, that is, how "wide" or "broad" it is.

The smoothed time series s_t is then given by:

$$\begin{aligned} s_t &= \sum_{i=1}^n a_i(t)x_t \\ &= \frac{\sum_{i=1}^n K\left(\frac{t-i}{b}\right)x_t}{\sum_{j=1}^n K\left(\frac{t-j}{b}\right)} \end{aligned}$$

The R function `ksmooth()` with the `normal` kernel argument provides exponential smoothing. The `bandwidth` parameter determines the "width" of the kernel by specifying the distance from the quartiles to the mean of the normal distribution function. Example smoothing results for various bandwidth values are shown in Figure 12.7.

```
# Apply gaussian kernel smoothing
ksmooth(time(soi), soi, kernel='normal', bandwidth=1)
```

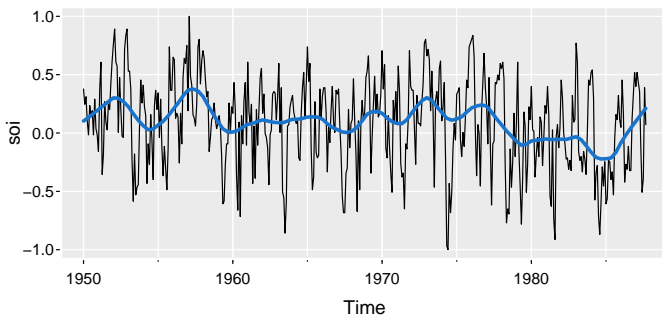


Figure 12.8: Lowess regression smoothing example

Lowess Regression

Lowess Regression (locally weighted scatterplot smoothing) combines a multiple regression model with a k-nearest-neighbour-based model. Each point on the smoothed time series is estimated by a weighted least squares regression over a local neighbourhood of f observations that are closest in time to the target point. The weights decrease with distance from the target observation, thereby providing robustness against outliers and yielding a smooth curve that closely follows the data.

The R function `lowess()` uses the `f` parameter for specifying the proportion of observations used for the regressions. An example result is shown in Figure 12.8.

```
# Apply lowess smoothing
lowess(soi, f=0.1)
```

Smoothing Splines

Smoothing splines are a method that fits a smooth, flexible “spline” function to the data. Smoothing splines balance the fit of the spline to the data against the smoothness of the spline curve and are essentially penalized polynomial regression models that fit the model:

$$m_t = \beta_0 + \beta_1 t + \beta_2 t^2 + \beta_3 t^3$$

by minimizing the loss function

$$\sum_{t=1}^n (x_t - m_t)^2 + \lambda \int \left(\frac{d^2 m}{dt^2} \right)^2 dt$$

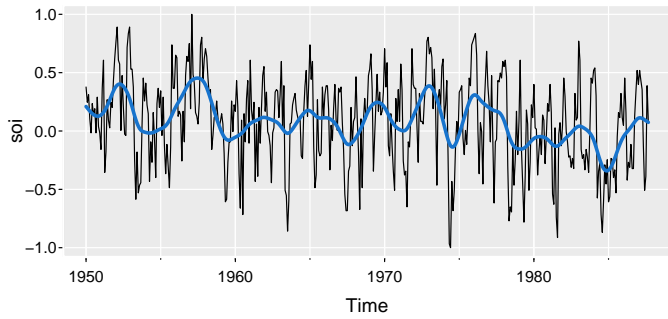


Figure 12.9: Smoothing spline example

The `smooth.spline()` function uses cubic splines, that is polynomials of degree 3. The smoothing parameter `spar` controls the regression penalty λ in the equation above and thereby the degree of smoothing. The result for this example is shown in Figure 12.9.

```
# Apply smoothing splines
smooth.spline(time(soi), soi, spar=0.5)
```

Hands-On Exercise

1. Generate 100 observations from the autoregression model $x_t = -.9x_{t-2} + w_t$ with $\sigma_w^2 = 1$
 - (a) Smooth the time series using a moving average filter $v_t = (x_t + x_{t-1} + x_{t-2} + x_{t-3})/4$, plot x_t as a line and superimpose v_t . Comment on the behaviour of x_t and how applying the moving average filter changes that behavior
 - (b) Smooth the time series using kernel smoothing, produce plots as above, and experiment with different kernel bandwidths. Comment on the behaviour of the smoothed series.
 - (c) Smooth the time series using Lowess, produce plots as above, and experiment with different values for the fraction of observations to include in each regression. How does the smoothed series change as you vary that fraction?
 - (d) Smooth the time series using smoothing splines, produce plots as above, and experiment with different values for the smoothing parameter that controls the regression penalty. How does the smoothed series change as you vary that parameter?
2. Generate 100 observations from the sinusoidal series $x_t = \cos(2\pi t/4)$ and add $N(0, 1)$ noise. Repeat the four smoothing exercises. Compare and contrast the results of these exercises. Which smoothing is more appropriate for which type of time series data?

12.5 Time Series Regression

Time series regression refers to using time series data in ordinary least squares regression. The focus is not necessarily on modeling data series over time or describing the future values of a time series as a function of earlier values, although lagged time series can certainly be used in time series regression. Instead, time series regression predicts the value of one time series from one or more other time series.

Consider an epidemiological example with three weekly time series, one expressing cardiovascular mortality ("cmort", the likelihood of dying of heart attack), another describing ambient temperature ("tempr") and a third one describing air pollution ("part"). Figure 12.10 shows these three time series superimposed in one graph. Visual inspection of the graph suggests that mortality may be correlated with temperature and air pollution.

Time series regression uses ordinary least squares (OLS) regression models where each series is a predictor variable. It essentially neglects the time aspect of the time series data. The following R code block shows examples of time series regression to predict or explain mortality. Note the use of the `time()` function extract the timestamps from the time series data.

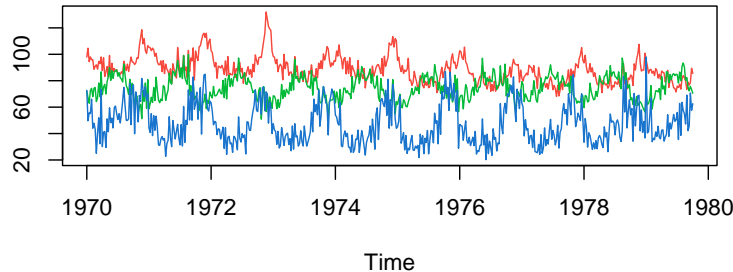


Figure 12.10: Three time series

```
# Use data from the astsa library
library(astsa)
# Plot the three time series
ts.plot(cmort, temp, part, col=2:4)
# Center the temperature variable
temp = temp - mean(temp)
# Square the temperature variable
temp.2 = temp^2
# Fit different linear models and provide summaries
summary(lm(cmort ~ time(cmort)))
summary(lm(cmort ~ time(cmort) + temp))
summary(lm(cmort ~ time(cmort) + temp + temp.2))
summary(lm(cmort ~ time(cmort) + temp + temp.2 + part))
```

Time series regression can also use lagged time series data, that is, data of the same series that is shifted backward in time. This is somewhat similar to the autoregressive models defined below. The following R code block lags the temperature by two weeks and by four weeks. It then uses `ts.intersect()` to combine the time series in a data frame for the times where they intersect. The data frame is then used in an OLS regression; different models could be fitted to identify the best explanation of mortality.

```
# Lag the temperature
temp.l.2 = lag(temp, 2)
temp.l.4 = lag(temp, 4)
# Intersect all time series to omit leading/trailing NA
temp.df <- ts.intersect(cmort, time(cmort), part,
                        temp, temp.2, temp.l.2, temp.l.4, dframe=TRUE)
# Fit the linear model including lagged temperature
summary(lm(cmort ~ time.cmort. + temp + temp.2 +
           temp.l.2 + temp.l.4 + part, data=temp.df))
```

12.6 Stationarity

The concept of *stationarity* is central to time series analysis. Stationarity means that the statistical characteristics of a time series do not change over time. That is, its mean, variance, and autocorrelation (the correlation of a time series with a lagged copy of itself) remain constant over time. Understanding and ensuring stationarity in a time series is important for the effective application of many statistical forecasting methods and models.

Stationary data with a constant mean and variance is more predictable and therefore easier to model. Changes in mean and variance can lead to forecasts that are biased or that degrade in accuracy over time. Stationarity ensures that the properties of the series used to generate forecasts will be similar in the future, which is crucial for planning and decision-making. If a time series is non-stationary, the behavior of the data could change over time, leading to models that are invalid or inaccurate when applied to future data points.

Statistical inference in time series analysis relies heavily on the assumption of stationarity. Many time series statistical models, including linear regression and ARMA models, are based on the assumption of stationarity. These models provide meaningful and reliable results only if the stationarity assumption is satisfied.

Strict stationarity is defined as the requirement that the probabilistic behaviour of every set of values of the series

$$\{x_{t1}, x_{t2}, \dots, x_{tk}\}$$

is identical to that of the set of values shifted by time h :

$$\{x_{t1+h}, x_{t2+h}, \dots, x_{tk+h}\}$$

That is,

$$\Pr\{x_{t1} \leq c_1, \dots, x_{tk} \leq c_k\} = \Pr\{x_{t1+h} \leq c_1, \dots, x_{tk+h} \leq c_k\}$$

Because strong stationarity is hard to test, a more commonly used and practical form of stationarity is *weak stationarity*, which requires only that the mean, variance, and the *autocovariance* (the covariance of the series with a lagged version itself) are constant over time. Most statistical tests and models assume weak stationarity. In summary, a weakly stationary time series is a finite variance process such that:

1. The mean and variance are constant and do not depend on time: $\mu_t = \mu$, $\sigma_t = \sigma$
2. The autocovariance γ depends on s and t only through their difference $h = |s - t|$.

Let $s = t + h$, then under the assumption of weak stationarity:

$$\begin{aligned} \gamma(s, t) &= \gamma(t + h, t) && \text{(because of condition 2)} \\ &= \text{cov}(x_{t+h}, x_t) && \text{(because of condition 1)} \\ &= \text{cov}(x_h, x_0) = \gamma(h) && \text{(autocovariance for lag } h\text{)} \end{aligned}$$

and

$$\rho(h) = \gamma(h)/\gamma(0) \quad (\text{autocorrelation for lag } h)$$

The *autocovariance* and *autocorrelations* are measures of dependence of the time series on lagged versions of itself. For a weakly stationary time series, the theoretical autocovariance for a lag h is defined as the covariance between two points $t, t + h$ on time series x

$$\gamma(h) = \text{cov}(x, x_{t+h}) = E[(x_t - \mu)(x_{t+h} - \mu)]$$

Note that this definition implies weak stationarity because a constant term for the mean μ is used in the expectation on the right-hand side.

A large autocovariance indicates a "smooth" time series, as each future value is strongly dependent on the previous value(s). In contrast, a small autocovariance indicates the "choppy" time series, as there is less dependence on prior values and values of the time series are less constrained and allowed to vary more.

The sample autocovariance that can be estimated from a finite sample for lag h is defined as

$$\hat{\gamma}(h) = \frac{1}{n} \sum_{t=1}^{n-h} (x_t - \bar{x})(x_{t+h} - \bar{x})$$

The *autocorrelation function* (ACF) for lag h is defined as usual as the autocovariance divided by the root of the product of the variances of the two time series:

$$\rho_x(h) = \frac{\gamma(t+h, t)}{\sqrt{\gamma(t+h, t+h)\gamma(t, t)}} = \frac{\gamma(h)}{\gamma(0)} \quad (\text{weak stationarity})$$

Note that this assumes weak stationarity. The time series properties at any time t are the same as at time 0 so that the above equation can be reduced to the right-most term.

Similar to the sample autocovariance, the sample ACF for lag h is defined as

$$\hat{\rho}_x(h) = \frac{\hat{\gamma}(h)}{\sqrt{\hat{\gamma}(h)\hat{\gamma}(0)}} = \frac{\hat{\gamma}(h)}{\hat{\gamma}(0)} \quad (\text{weak stationarity})$$

where the last step again assumes weak stationarity.

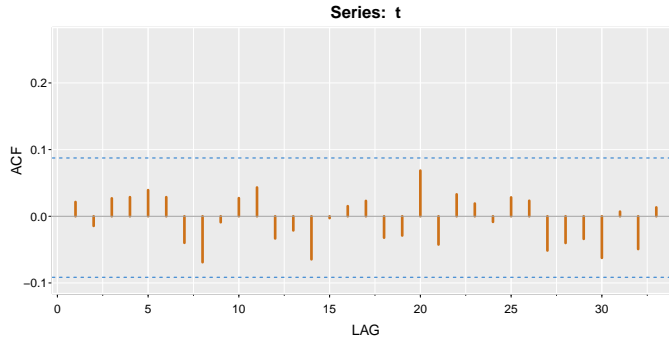


Figure 12.11: ACF of Gaussian white noise

To test whether the ACF of any sequence for lag h is statistically different from 0, note that the large-sample distribution of $\hat{\rho}_x(h)$ is normal with mean 0 and standard deviation

$$\sigma_{\hat{\rho}_x} = 1/\sqrt{n}$$

if the generating processes is independent white noise. Hence, the approximate 95% confidence interval on the ACF is

$$-\frac{1}{n} \pm \frac{2}{\sqrt{n}}$$

If the sample ACF of n values of a time series for a given lag exceeds the lower or upper bounds of the confidence interval, the ACF is statistically significantly different from 0, and the time series is unlikely to be white noise.

The following R code block illustrates the autocorrelation function using the standard `cor()` function to compute the correlations at different lags and the `acf1()` function of the `astsa` library that will automatically lag the time series and output and optionally plot the ACF values at different lags, creating a plot as in Figure 12.11.

```
library(astsa)
# Create Gaussian white noise
t <- ts(rnorm(500))
# The hard way:
cor(ts.intersect(t, lag(t,1), dframe=T))
cor(ts.intersect(t, lag(t,2), dframe=T))
# etc.
# The easy way:
# Without plot
acf <- acf1(t, plot=FALSE)
# With plot
acf1(t, gg=T, col=7, lwd=3)
```

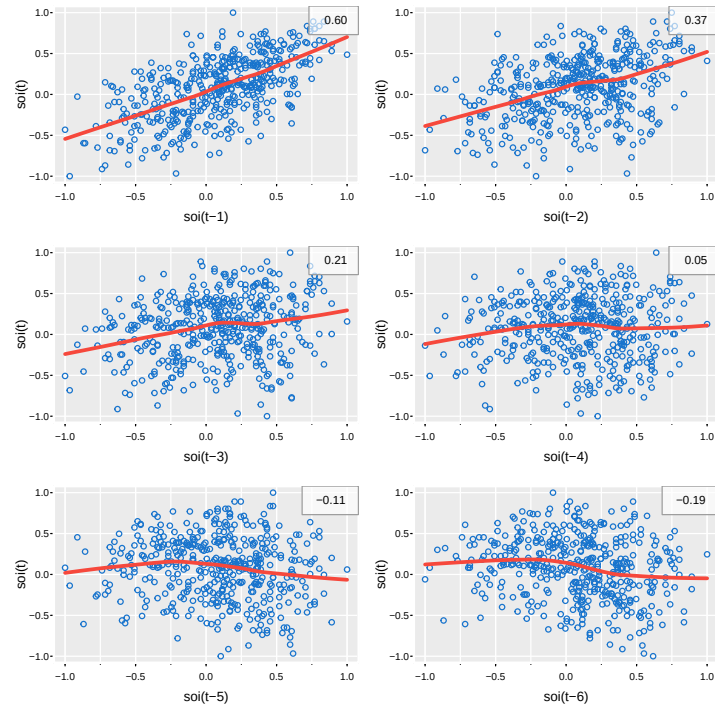


Figure 12.12: Autocorrelations at six different lags

The following example in R uses the `soi` data set and the `lag1.plot()` from the `astsa` library to provide also a graphical display of the autocorrelations at various lags, as shown in Figure 12.12.

```
library(astsa)
# Compute and plot the ACF for different lags
acf1(soi, gg=T, co=3, lwd=2)
# Scatterplot of original versus or lags up to 6, with ACF values
lag1.plot(soi, max.lag = 6, gg=T, col=4, lwl=3)
```

The *partial autocorrelation function* (PACF) of a time series is a measure of the correlation between observations at two points in time, accounting for the correlations of the observations at all shorter intervals. Essentially, it reflects the direct effect of past data points on the future data point, after removing the effects of intermediate data points. PACF can be thought of as the correlation between a variable and its lag h that is not explained by correlations at all lower-order lags. It is formally defined as the correlation between x_{t+h} and x_t with the linear dependence of $\{x_{t+1}, \dots, x_{t+h-1}\}$ on each removed:

$$\phi_{hh} = \begin{cases} \rho(1) & h = 1 \\ \text{corr}(x_{t+h} - \hat{x}_{t+h}, x_t - \hat{x}_t) & h \geq 2 \end{cases}$$

The following R code block illustrates the use of the partial autocorrelation function of a time series, first using the standard `cor()` function for a lag of 3 and then the `acf1()` function of the `astsa` library that automatically computes the PACF for different lags.

```
t <- ts(rnorm(500))
# The hard way
# Shift the series to create lagged versions
t1 <- lag(t, 1)
t2 <- lag(t, 2)
t3 <- lag(t, 3)
data <- ts.intersect(t, t1, t2, t3, dframe=T)

# Using linear models to adjust for intervening lags
model_lag1 <- lm(t ~ t1 + t2, data)
model_lag2 <- lm(t1 ~ t2, data)
# Residuals for lag 3
residuals_lag1 <- residuals(model_lag1)
residuals_lag2 <- residuals(model_lag2)
final_model <- lm(residuals_lag1 ~ residuals_lag2)
# Correlation between residuals and lag 3 data
pacf_lag3 <- cor(residuals(final_model), data$t3)

# The easy way
acf1(t, plot=F, pacf=T)
```

12.7 Dealing with Non-Stationarity

When a time series is non-stationary, it can often be transformed into a stationary series through techniques such as logarithmic or square root transformations, detrending, and differencing. These transformations can stabilize the mean and reduce variance dependency over time.

Transformations

Popular *time series transformations* are the log transformation, the square root transformation and the Box-Cox power transformation, defined as follows:

$y_t = \log x_t$	Log transformation
$y_t = \sqrt{x_t}$	Square root transformation
$y_t = \begin{cases} (x_t^\lambda - 1)/\lambda & \lambda \neq 0 \\ \log x_t & \lambda = 0 \end{cases}$	Box-Cox power transformation

Detrending

Detrending a time series involves removing the trend component from the data, thereby isolating the non-trend components such as seasonality and irregular fluctuations. This is particularly useful in time series analysis because many statistical methods assume stationarity (constant mean and variance), and a trend violates these assumptions.

A common detrending method is to fit a regression model to the trend component and then subtract the fitted values, that is, the trend, from the original series. Linear regression is widely used for linear trends, but polynomial or more complex models can be fitted depending on the nature of the trend.

For example, assume that

$$x_t = \mu_t + y_t$$

where μ_t is the trend and y_t a stationary series. Then detrending comprises the following two steps:

1. Estimate trend, e.g. with a linear model such as $\mu_t = \beta_0 + \beta_1 t$
2. Work with residuals, e.g. $\hat{y}_t = x_t - \hat{\mu}_t = x_t - \hat{\beta}_0 - \hat{\beta}_1 t$

The following R code block shows how to detrend a time series using linear regression, producing the graphs shown in Figure 12.13.

```
# Simulate a time series with a linear trend
t <- ts(1:100 + rnorm(100) * 10)

# Fit a linear model to the time series
trend_model <- lm(t ~ time(t))
# Calculate detrended series by subtracting the estimated trend
detrended_series <- residuals(trend_model)

# Plot original and detrended
par(mfrow=c(2,1))
tsplot(t, type="l", main="original", col=3, gg=T)
tsplot(detrended_series, type="l", main="detrend", col=2, gg=T)
```



Figure 12.13: Time series and detrended time series

Differencing

Differencing involves computing the differences between consecutive observations in the original time series. The primary goal of differencing is to remove trends and seasonality in order to stabilize the mean of the time series by reducing changes in the level of a time series over time. Assume again that

$$x_t = \mu_t + y_t$$

where μ_t is the trend and y_t a stationary series. Differencing models the trend stochastically as a random walk with drift:

$$\mu_t = \delta + \mu_{t-1} + w_t$$

where w_t is white noise. Differencing then yields

$$\begin{aligned} x_t - x_{t-1} &= (\mu_t + y_t) - (\mu_{t-1} + y_{t-1}) \\ &= \delta + w_t + y_t - y_{t-1} \end{aligned}$$

which is stationary.

As seen above, the first difference can remove a linear trend. However, sometimes the first difference is not enough to achieve stationarity. In such cases, the second

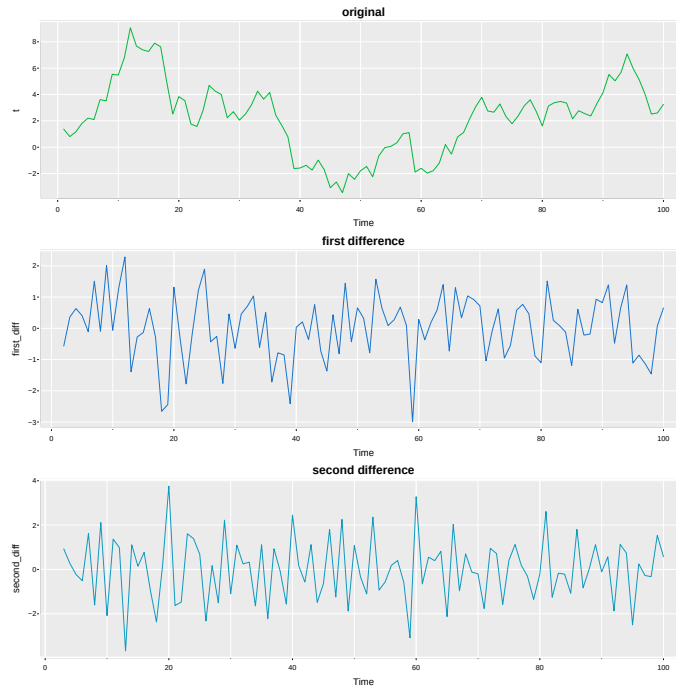


Figure 12.14: Original, first and second differences of a simulated time series

difference can be used to remove a quadratic trend and higher-order differences can be computed if the series still shows non-stationary behavior after the second differencing.

The following R code shows the effect of differencing on a simulated time series. Differencing uses the `diff()` function. The resulting plots are shown in Figure 12.14.

```
# Simulating a time series with trend
t <- ts(cumsum(rnorm(100))) # Cumulative sum of normal deviations

par(mfrow=c(3,1))
tsplot(t, type="l", main="original", col=3, gg=T)
# First differencing
tsplot(diff(t, differences = 1), type="l",
       main="first difference", col=4, gg=T)
# Second differencing
tsplot(diff(t, differences = 2), type="l",
       main="second difference", col=5, gg=T)
```

To see illustrate the effects of detrending and differencing on the ACF for a real time series, consider the chicken price data set `chicken` in the `astsa` library. Figure 12.15 shows the ACF for the original, the detrended, and the differenced series (first and second differences). While the original time series is clearly non-stationary with large

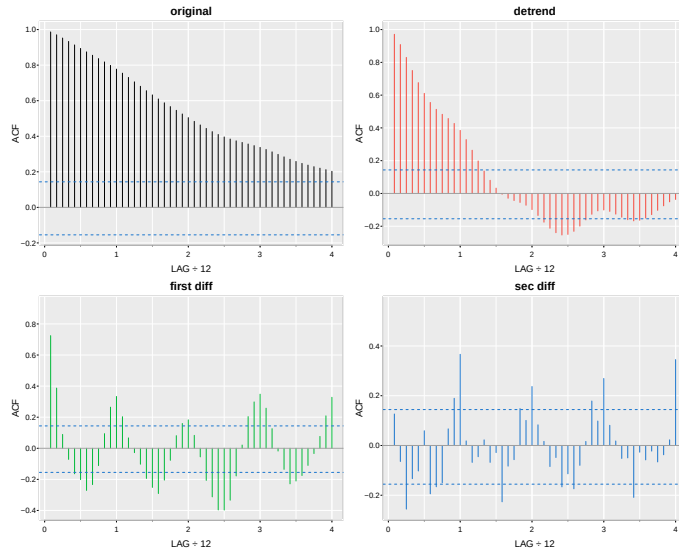


Figure 12.15: ACF for detrended and differenced time series

ACF values (top left panel), the detrended series improves this somewhat, but still shows large ACF (top right panel). First differencing reduces the ACF values and shows a cyclical trend with a cycle of 6 months, with significant ACF values (bottom left panel). The second difference in the bottom right panel of Figure 12.15 still shows significant ACF values at the 6 month and 12 month lags but non-significant ACF for most other lags.

```
acfl(chicken, max.lag=48, main="original", col=1, gg=T)
acfl(resid(fit), max.lag=48, main="detrend", col=2, gg=T)
acfl(diff(chicken), max.lag=48, main="first diff", col=3, gg=T)
acfl(diff(chicken, differences=2), max.lag=48,
     main="sec diff", col=4, gg=T)
```

Hands-On Exercises

1. Extend the mortality, temperature and pollution/particulate model by adding another component to the regression that accounts to the particulate four weeks prior; that is, add the lagged pressure P_{t-4} to the regression.
2. Draw a scatterplot matrix of mortality M_t , temperate T_t , pressure P_t and lagged pressure P_{t-4} , then calculate the pairwise correlations between them. Compare the relationship between M_t and P_t versus M_t and P_{t-4} .

Source: Shumway & Stoffer, Chapter 2

Hands-On Exercises

1. Detrend the `soi` time series data by fitting a regression of S_t on time t . Is there a significant trend in the surface pressure?
2. Use two different smoothing techniques to estimate the trend in the global temperature series `gtemp_both` in the `astsa` library.

Source: Shumway & Stoffer, Chapter 2

Hands-On Exercise

Consider the two weekly time series `oil` and `gas` in the `astsa` library. The oil series is in dollars per barrel, while the gas series in in cents per gallon.

1. Plot the data on the same graph. Do you believe the series are stationary?
2. Apply the transformation $y_t = \nabla \log x_t$ to the data for both series
3. Plot the transformed series on the same graph, and calculate the ACFs for both series
4. Plot the CCF of the transformed series and comment.

Source: Shumway & Stoffer, Chapter 2

12.8 ARIMA Models

ARIMA models, which stands for Autoregressive Integrated Moving Average, are a type of statistical models for analyzing and forecasting time series data. ARIMA is particularly suited to time series data that show non-stationarities, such as trends and seasonal patterns, and it has become a standard tool in econometrics, finance, and other fields.

ARIMA models can be divided into the following model classes:

- **AR:** pure AutoRegressive models
- **MA:** pure Moving average models
- **ARMA:** model with AutoRegressive and Moving-Average terms
- **ARIMA:** AutoRegressive Integrated Moving-Average models (involves differencing for non-stationary time series with trend)

To simplify working with ARIMA models, the difference operator ∇ is defined as:

$$\nabla x_t = x_t - x_{t-1}$$

Building on this definition, the *Backshift operator* or *Lag Operator* B is defined as:

$$\begin{aligned} B x_t &= x_{t-1} \\ B^k x_t &= x_{t-k} \\ \nabla x_t &= (1 - B)x_t \\ \nabla^2 x_t &= (1 - B)^2 x_t \\ &= (1 - 2B + B^2)x_t \\ &= x_t - 2x_{t-1} + x_{t-2} \\ \nabla^d &= (1 - B)^d \end{aligned}$$

An *autoregressive model* of order p , denoted by AR(p), models the current value of a time series as a linear combination of previous values. The number of lagged observations used in the model is denoted by the order p . The AR model captures the regression of the time series on its previous values, which indicates persistence, or memory, within the series. It is defined as:

$$x_t = \phi_1 x_{t-1} + \phi_2 x_{t-2} + \cdots + \phi_p x_{t-p} + w_t$$

where w_t is white noise and the ϕ_i are model parameters.²

The *autoregressive operator* $\phi(B)$ is defined using the backshift operator as:

$$\begin{aligned} \phi(B) &= 1 - \phi_1 B - \phi_2 B^2 - \cdots - \phi_p B^p \\ &= \left(1 - \sum_{j=1}^p \phi_j B^j \right) \end{aligned}$$

²In contrast to an "ordinary" regression model, the x_i are random effects, not fixed, because each x_i has an associated error term w_t . This means that AR or ARIMA models in general are not estimated using OLS because the OLS assumptions are not met. Instead, AR and ARIMA models are estimated using maximum-likelihood or other methods.

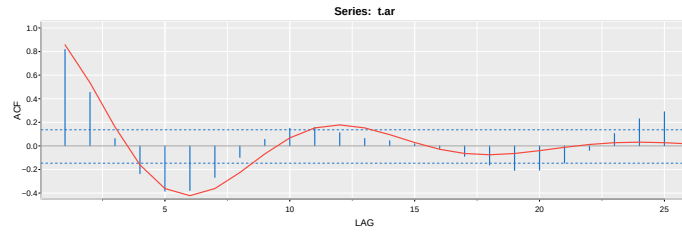


Figure 12.16: Simulated (blue) and theoretical (red) ACF of an AR(2) model

so that the AR(p) model becomes:

$$\phi(B)x_t = w_t$$

The theoretical ACF of a given AR(p) model can be calculated analytically. In R, the `ARMAacf()` function can be used for this by specifying the autoregressive coefficients ϕ . The following R code block simulates 200 observations of an AR(2) time series and plots the simulated (blue) versus theoretical (red) ACF values, shown in Figure 12.16. The theoretical values can be used to determine whether a specific time series conforms to a particular AR(p) model. The ACF of an AR(p) model is characterized by a slow decline of its values past a lag of p , as shown in Figure 12.16.

```
# Theoretical ACF of an AR(2) model
ARMAacf(ar=c(1.5, -.75), lag.max=10)
# Simulate an ARIMA(2,0,0) model with those AR coefficients
t.ar = arima.sim(list(ar=c(1.5, -.75)), n=200)
# Compute and plot the ACF of the simulated series
acf1(t.ar, max.lag=25, qq=T, lwd=2, col=4)
# Add the theoretical values for comparison
lines(ARMAacf(ar=c(1.5, -.75), lag.max=26)[-1], lwd=2, col=2)
```

A *moving average model* or order q , denoted by MA(q), models the current value of the series as a linear combination of past forecast errors, which are computed as differences between past values and their respective forecasts. The parameter q specifies the number of lagged forecast errors in the prediction equation. The MA model is useful for capturing "shock errors" in the model, providing a way to allow the model to adapt to sudden changes in the series. It is defined as:

$$x_t = w_t + \theta_1 w_{t-1} + \theta_2 w_{t-2} + \cdots + \theta_q w_{t-q}$$

where w_t are Gaussian errors and θ_i are model parameters.

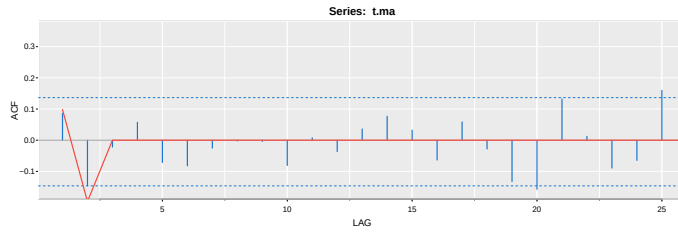


Figure 12.17: Simulated (blue) and theoretical (red) ACF of an MA(2) model

The *moving average operator* $\theta(B)$ is defined using the backshift operator as:

$$\begin{aligned}\theta(B) &= 1 + \theta_1 B + \theta_2 B^2 + \cdots + \theta_q B^q \\ &= \left(1 + \sum_{j=1}^q \theta_j B^j \right)\end{aligned}$$

so that the MA(q) model becomes:

$$x_t = \theta(B)w_t$$

The theoretical ACF of a given MA(q) model can be calculated analytically. In R, the `ARMAacf()` function can be used for this, by specifying the moving average coefficients θ . Similar to the previous example, the following R code simulates 200 observations of an MA(2) model and plots the simulated (blue) versus theoretical (red) ACF values, shown in Figure 12.17. In contrast to an AR(2) model, the ACF does not gradually diminish, but becomes 0 after lag q . The simulated values in Figure 12.17 confirm this as they are largely statistically non-significant past a lag of 2.

```
# Theoretical ACF of an MA(2) model
ARMAacf(ma=c(1.5, -.75), lag.max=10)
# Simulate an ARIMA(0,0,2) model with those MA coefficients
t.ma = arima.sim(list(ma=c(1.5, -.75)), n=200)
# Compute and plot the ACF of the simulated series
acf1(t.ma, gg=T, lwd=2, col=4)
# Add the theoretical values for comparison
lines(ARMAacf(ma=c(1.5, -.75), lag.max=26)[-1], lwd=2, col=2)
```

An *autoregressive moving-average model* of order (p, q) , denoted by $ARMA(p, q)$, combines both autoregressive and moving-average terms in the same model:

$$x_t = \alpha + \phi_1 x_{t-1} + \cdots + \phi_p x_{t-p} + w_t + \theta_1 w_{t-1} + \cdots + \theta_q w_{t-q}$$

Using the AR and MA operators defined above, this model can be written as:

$$\phi(B)x_t = \theta(B)w_t$$

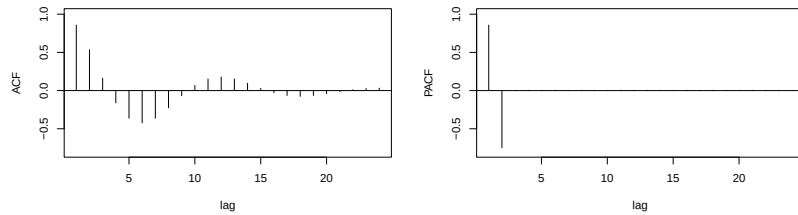


Figure 12.18: ACF and PACF of an AR(2) model

It turns out that every ARMA model has an equivalent MA only model. However, this equivalent MA model in theory has an infinite number of MA terms. In practice, a reasonable approximation can be achieved by retaining a limited number of MA terms.

Moreover, many ARMA models (the class of invertible ones) have an equivalent AR models. Again, this equivalent model has an infinite number of AR terms and again, in practice, reasonable approximations can be achieved by retaining a limited number of AR terms.

Equivalent models can be found using the `ARMAtoMA()` and `ARMAtoAR()` functions in the `astsa` library, which return the MA and AR coefficients of the equivalent models, as shown in the following R code example:

```
library(astsa)
# MA coefficients of equivalent MA models
ARMAtoMA(ar = c(1.5, -.75), lag.max=10)
ARMAtoMA(ar = c(-.5), ma = c(-.9), lag.max=10)
# AR coefficients of equivalent AR models
ARMAtoAR(ma = c(1.5, -.75), lag.max=10)
ARMAtoAR(ar = c(-.5), ma = c(-.9), lag.max=10)
```

As shown in Figures 12.16 and 12.17, the ACF of AR and MA models behave differently. Similarly, the PACF behaves differently for the two types of models. Figure 12.18 shows the ACF and the PACF for an AR(2) model. While the ACF diminishes gradually, the PACF is zero immediately after lag 2. These properties of the ACF and PACF can be used to select a suitable statistical model to fit a given time series, as shown in Table 12.1. When a AR gradually diminishes and the PACF cuts off suddenly after a lag p , this is an indication that an AR(p) model is suitable. Conversely, when the ACF cuts off suddenly after lag q and the PACF diminishes gradually, this is an indication for an MA(q) model. When neither ACF nor PACF cut off suddenly, a mixed ARMA(p, q) model should be fitted.

A full *autoregressive integrated moving average* ARIMA(p, d, q) model adds a differencing term to the ARMA(p, q) model to achieve weak stationarity of the time series.

	AR(p)	MA(q)	ARMA (p, q)
ACF	Tails off	Cuts off after lag q	Tails off
PACF	Cuts off after lag p	Tails off	Tails off

Source: Shumway&Stoffer, Table 3.1

Table 12.1: Properties of the ACF and PACF for AR and MA models

The AR operator can be factorized by $(1 - B)$, so that:

$$\begin{aligned}\phi(B) &= \left(1 - \sum_{j=1}^{p'} \phi_j B^j\right) \\ &= \left(1 - \sum_{j=1}^{p'-d} \phi_j B^j\right) (1 - B)^d\end{aligned}$$

With $p = p' - d$, the ARIMA(p,d,q) model is then:

$$\left(1 - \sum_{j=1}^p \phi_j B^j\right) (1 - B)^d x_t = \left(1 + \sum_{j=1}^q \theta_j B^j\right) w_t$$

This can be generalized to:

$$\left(1 - \sum_{j=1}^p \phi_j B^j\right) (1 - B)^d x_t = \delta + \left(1 + \sum_{j=1}^q \theta_j B^j\right) w_t$$

12.9 Fitting an ARIMA Model

Fitting an ARIMA model to time series data involves the following steps, from initial data analysis and transformation to final model selection:

1. Plot the data
2. Possibly transform the data
3. Assess stationarity
4. Possibly difference the data
5. Identify the dependence orders (p, q) of the model
6. Estimate parameters
7. Model diagnostics

8. Model selection

Plotting the data is useful as an initial visual assessment of stationarity, trends, or seasonality. A number of transformations have been discussed earlier that may be useful to "stabilize" a time series. If the series after transformation is still not stationary, differencing can remove trends and seasonal components. Determining the order of differencing needed to achieve stationarity is often done by trial and error, reassessing stationarity after each difference. A slow decay in the sample ACF $\hat{\rho}(h)$ typically indicates a need for differencing. However, over-differencing can introduce dependence where none actually exists. Typically, differencing should be done in small steps, beginning with a first difference, and then repeatedly checked with the ACF.

Identifying the initial ARMA order p and q should be done based on the ACF and PACF functions, using Table 12.1 as a basis. Often, multiple model may need to be tried, altering the AR and MA orders p and q in small steps.

The following R code example uses a quarterly time series of the US gross national product (gnp) from the `astsa` library as an example. The `acf2()` function of the `astsa` library produces simultaneous plots of ACF and PACF for ease-of-use. The time series is then log-transformed and differenced once. The results are shown in Figure 12.19.

```
# Plot data
plot(gnp)
# Plot ACF
acf2(gnp, 50)
# Log transform, and first order differencing
gnpgr = diff(log(gnp))
# Plot transformed and differenced data
plot(gnpgr)
# Plot ACF of transformed and differenced data
acf2(gnpgr, 24)
```

Note how the ACF of the original series diminishes very gradually, indicating the need for differencing. The sample ACF of the transformed and differenced series shows a gradual decline after a lag of 2, while the sample PACF of the transformed and differenced series cuts off to non-significance after a lag of 1. Together, this indicates that the time series may be appropriately modeled using an AR(1) model or an MA(2) model. Converting the AR(1) model to an equivalent MA model shows that the two initial models are approximately equivalent. Note that the following R code block uses the `sarima()` function from the `astsa` library because this function also produces the diagnostic plots shown in Figures 12.20 and 12.21. This function can fit ARIMA models, as in the following example, but can also fit seasonal ARIMA, or SARIMA, models.

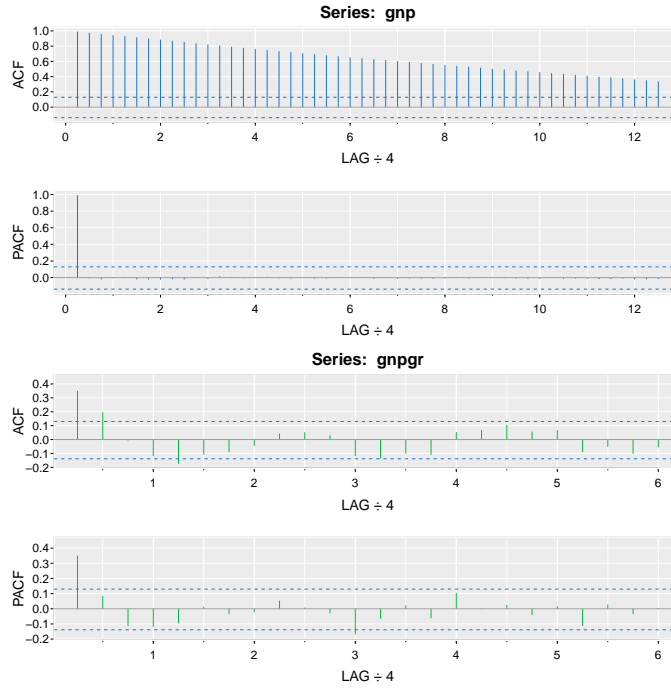


Figure 12.19: ACF and PACF of the original time series (top) and of the log-transformed and differenced series (bottom)

```
# Fit an AR(1) model
sarima(gnpgr, 1, 0, 0)
# Fit an MA(2) model
sarima(gnpgr, 0, 0, 2)
# Models are roughly equivalent
ARMAToMA(ar=0.35, ma=0, 10)
```

ARIMA model diagnostics focus on the residuals of the fitted model and typically assess the following criteria:

- Standardized residuals should be Gaussian ($\mu = 0$, $sd = 1$)
- Residuals should not be autocorrelated
- Residual ACF should be Gaussian with $\mu = 0$ and $sd = 1/\sqrt{n}$
- Ljung-Box statistic Q of the error ACF $\hat{\rho}_e$ for different maximum lags H should be larger than the $1 - \alpha$ quantile of the χ^2_{H-p-q} distribution (i.e. the test statistic

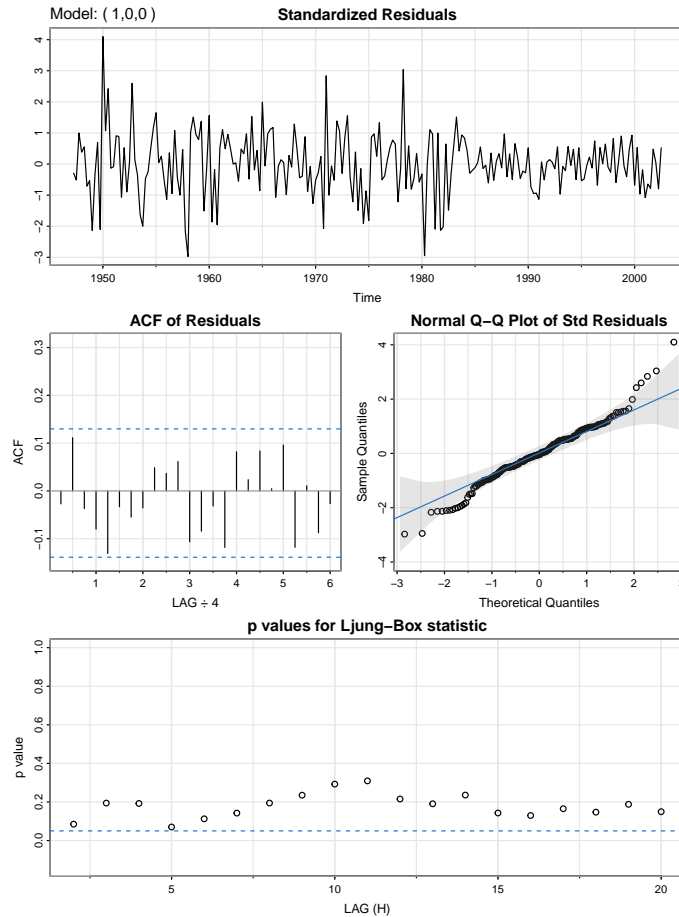


Figure 12.20: Diagnostics for an AR(1) model fitted to the GNP time series

is not statistically significantly different from 0)

$$Q = n(n + 2) \sum_{h=1}^H \frac{\hat{\rho}_e^2(h)}{n - h}$$

The diagnostic plots in Figure 12.20 and 12.21 show the residuals in the top panels. There are no visible trends or regularities and a few large outliers, but these can be expected from a Gaussian distribution. The ACF of the residuals in the middle left panel show that they are not autocorrelated and the QQ plot of residuals shows some deviations from a linear diagonal in the bottom and top portions, indicating that the model over- and under-estimates extreme values. The bottom panel in each figure shows the probability values (p-values) for the Ljung Box statistics and the $1 - \alpha$ dashed horizontal line. For both models, the p-values of the Ljung Box statistic are

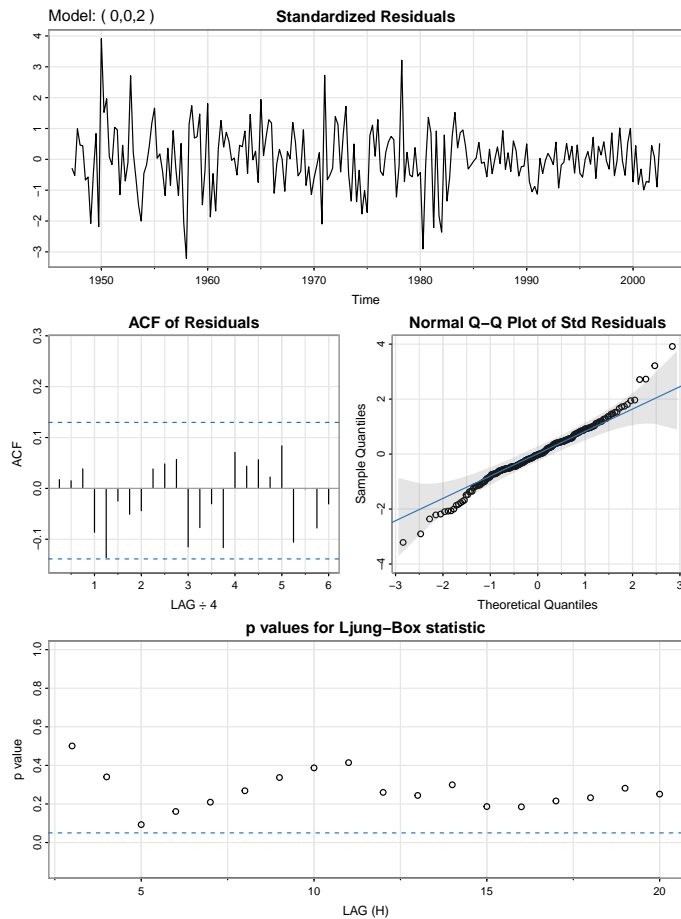


Figure 12.21: Diagnostics for an MA(2) model fitted to the GNP time series

above the horizontal line, that is, they are not significantly different. In summary, both models show good fit to the data.

Because fitting a model typically uses maximum-likelihood estimation (MLE), the model choice is often based on information criteria that are based on the log-likelihood L of the model. Because more complex models naturally achieve a better fit to the training data, that is, they have a smaller bias, the log-likelihood is adjusted (penalized) for model complexity, that is, the number of parameters k , and is also adjusted for sample size n . All information-theoretic criteria express a *relative* quality of fit with *smaller values being better*. There are no absolute cut-off values that would indicate a well-fitting model.

$$\text{AIC} = -2 \log L + 2k \quad \text{Akaike Information Criterion}$$

$$\text{AICc} = \text{AIC} + \frac{2k(k+1)}{n-k-1} \quad \text{Akaike Information Criterion, corrected}$$

$$\text{BIC} = -2 \log L + k \log n \quad \text{Bayesian Information Criterion}$$

The R output of the `sarima()` function shows very similar model fit values:

```
> sarima(gnpgr, 1, 0, 0)
AIC = -6.44694  AICc = -6.446693  BIC = -6.400958
```

```
> sarima(gnpgr, 0, 0, 2)
AIC = -6.450133  AICc = -6.449637  BIC = -6.388823
```

Once the analyst has selected the final model and is satisfied that it fits well, future values of the time series can be forecasted from the fitted model. An important property of ARIMA predictions is that they quickly settle to the mean, with a constant prediction error, reflecting the stationarity of the differenced and transformed time series.

The following R code example shows forecasting from using the `sarima.for()` function in the `astsa` library. The results are shown visually in Figure 12.22, where the prediction error is indicated by the gray shading.

```
forecasts <- sarima.for(gnpgr, n.ahead=10, p=1, d=0, q=0)
```

12.10 GARCH Models

General Autoregressive Conditional Heteroscedasticity (GARCH) models are a family of time series models that are used to estimate the volatility and conditional variance of time series data, particularly of financial time series that exhibit time-varying volatility and volatility clustering. GARCH models are fundamental in the field of financial econometrics for modeling financial time series data.

GARCH models predict the current variance (that is, the volatility or variability, not the actual values) as a function of past squared "innovations" (which represent unexpected shocks or news in the data) and past conditional variances. In other words, the variance at any time depends on the information available up to the previous period. GARCH models are particularly effective at modeling volatility clustering, a phenomenon common in financial time series where high-volatility events tend to cluster together.

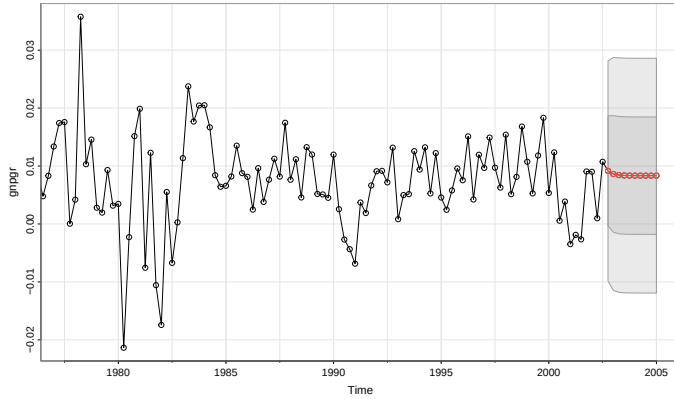


Figure 12.22: Forecasting from an ARIMA(1,0,0) model with estimated prediction errors

GARCH models are extensively used in risk management, asset pricing, and financial forecasting. They help in estimating the volatility of asset returns for pricing derivatives, calculating the value at risk for risk management, or forecasting volatility for portfolio optimization.

An ARCH model considers a series of "returns", which are defined as deviations from the prior value:

$$r_t = \frac{x_t - x_{t-1}}{x_{t-1}} \quad (\text{"Return"})$$

The series of returns is modelled as the product of a stochastic component ϵ_t and a time-dependent standard deviation σ_t

$$r_t = \sigma_t \epsilon_t$$

The ARCH model considers the time-dependent variance σ_t^2 at time t as a function of the previous returns. For example, in the ARCH(1) model the variance σ_t^2 at time t depends on the immediately prior squared return:

$$\sigma_t^2 = \alpha_0 + \alpha_1 r_{t-1}^2$$

where ϵ_t is Gaussian.

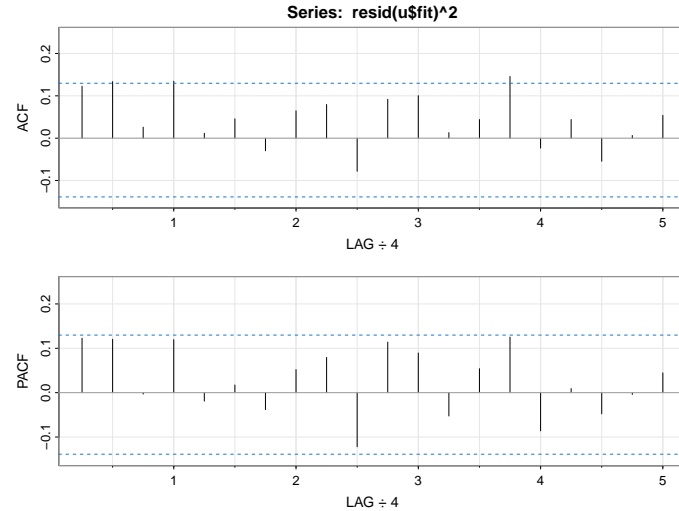


Figure 12.23: Squared residuals after fitting an AR(1)+ARCH(1) model to the US GNP time series data

The general ARCH(q) model of order q is defined as follows. Again, the variance depends on the prior squared returns:

$$\begin{aligned}\sigma_t^2 &= \alpha_0 + \alpha_1 r_{t-1}^2 + \alpha_2 r_{t-2}^2 + \dots + \alpha_q r_{t-q}^2 \\ &= \alpha_0 + \sum_{i=1}^q \alpha_i r_{t-i}^2\end{aligned}$$

ARCH models can be combined with ARIMA models so that the ARCH model describes the error term. A simple example is an AR(1) model with ARCH(1) error terms:

$$x_t = \phi_0 + \phi_1 x_{t-1} + \sigma_t \epsilon_t \quad \text{where} \quad \sigma_t = \alpha_0 + \alpha_1 x_{t-1}^2$$

As an example, consider the US gross national product quarterly time series from the `astsa` library. An initial AR(1) model shows that the squared residuals have some dependence. This dependence can be accounted for by explicitly modeling the residuals as ARCH(1). The `fGarch` library for R provides the `garchFit()` function to these kinds of models. The squared residuals of the final model are shown in Figure 12.23.

```

library(astsa)
# Fit an AR(1) model to the differenced, log-transformed series
u = sarima(diff(log(gnp)), 1, 0, 0)
# Examine the squared residuals
acf2(resid(u$fit)^2, 20)

library(fGarch)
# Fit an AR(1) + ARCH(1) model to the differenced, log-transformed
# series and show the summary
summary(garchFit(~arma(1,0)+garch(1,0), diff(log(gnp))))

```

An extension to ARCH is to model the variance not only as a function of previous returns, but also as a function of the p prior variances. In other words, the variance is modelled as an autoregressive model in addition to the ARCH(q) terms. This leads to a Generalized ARCH model, that is, a GARCH(p, q) model:

$$\begin{aligned}\sigma_t^2 &= \omega + \alpha_1 r_{t-1}^2 + \cdots + \alpha_q r_{t-q}^2 \\ &\quad + \beta_1 \sigma_{t-1}^2 + \cdots + \beta_p \sigma_{t-p}^2 \\ &= \omega + \sum_{j=1}^q \alpha_j r_{t-j}^2 + \sum_{j=1}^p \beta_j \sigma_{t-j}^2\end{aligned}$$

The following R example models the Dow Jones Industrial Average stock market time series values (data set `djiar` in library `astsa`) using an AR(1)+GARCH(1,1) model. Parameter estimates are shown below. Various diagnostic plots are available using the `plot()` function for the resulting `garchFit` object and are shown in Figure 12.24.

```

library(zoo)
library(fGarch)
# Log transform
djiar = diff(log(djia$Close))[-1]
# Fit an AR(1) + GARCH(1,1) model
djia.g <- garchFit(~arma(1,0)+garch(1,1), data=djiar)
# Show summary information
summary(djia.g)
# Different plots available
par(mfrow=c(5,2))
plot(djia.g, which=1:10)

```

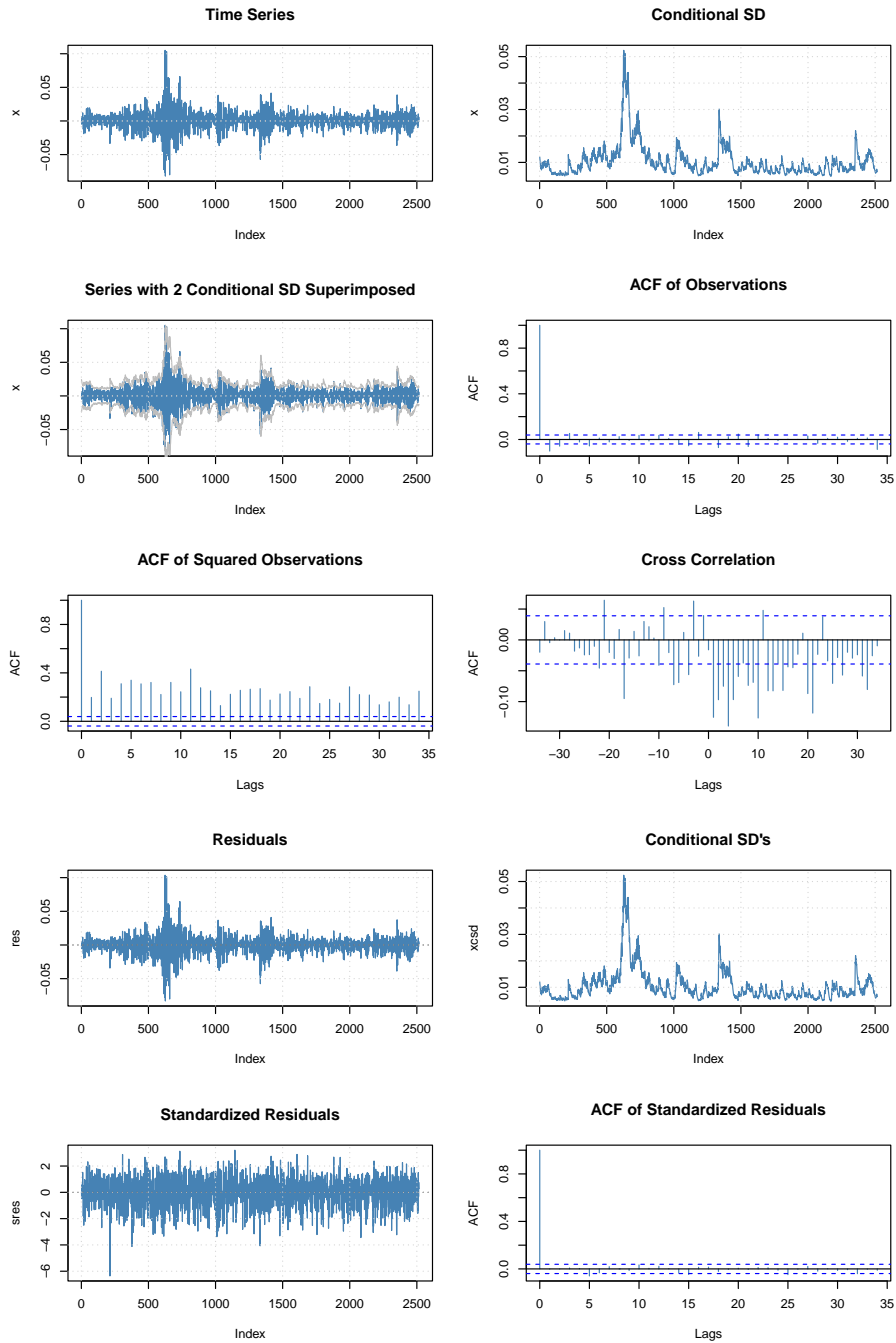


Figure 12.24: Diagnostic plots for a GARCH model

	Estimate	Std. Error	t value	Pr(> t)	
mu	8.585e-04	1.470e-04	5.842	5.16e-09	***
ar1	-5.532e-02	2.023e-02	-2.735	0.006238	**
omega	1.610e-06	4.459e-07	3.611	0.000305	***
alpha1	1.244e-01	1.660e-02	7.496	6.55e-14	***
beta1	8.700e-01	1.526e-02	57.022	< 2e-16	***
shape	5.979e+00	7.917e-01	7.551	4.31e-14	***
<hr/>					

Log Likelihood:					
8249.619 normalized: 3.27756					

Appendix – Basic Time Series Functions in R

base or stats	
filter	Filters time series, through moving averages or autoregression
lag	Creates a lagged version of a time series by shifting the time-base back
diff	Creates lagged differences
plot.ts	Plot a time series
ts.plot	Plot multiple time series
lag.plot	Scatterplot of lagged values
acf	ACF and plot
ccf	CCF and plot
time	Creates the vector or times at which a time series was sampled
cycle	Gives the positions in the cycle of each observation
frequency	Number of samples per unit time
ts.intersect	Bind time series together that have a common frequency. Restrict to time covered by all series
ts.union	Bind time series together that have a common frequency. Pad with NA if necessary
ar	Fit an autoregressive model
arima	Fit an ARIMA model
astsa	
tsplot	Plot a time series
acf1	ACF and plot
ccf2	CCF and plot
sarima	Fit seasonal ARIMA models (and nice diagnostic plots)
lag1.plot	Scatterplot of lagged values

12.11 Review Questions

Introduction

1. What is time series analysis and why is it important in various fields such as economics, finance, and natural sciences?
2. What are some essential preprocessing steps required before performing time series analysis?
3. Compare and contrast the time-domain approach and the frequency-domain approach in time series analysis. Which approach is particularly useful for forecasting and why?

Time Series Statistical Models

4. Explain the Moving Average (MA) model and describe how it uses past error terms to forecast future values. What are the assumptions about these error terms?
5. Discuss how the Moving Average (MA) model can be utilized to detect underlying patterns in a time series that exhibits random fluctuations. What limitations does this model have in handling trend and seasonality?
6. Explain the implications of choosing different window sizes (the number of terms included) in the Moving Average model. How does it affect the forecasts and smoothing?
7. Describe how the Autoregressive (AR) model differs from the MA model and provide an example of its application in economic or financial time series.
8. In the context of the AR model equation $x_t = x_{t-1} - 0.9x_{t-2} + w_t$, analyze the impact of changing the coefficient -0.9 to values closer to 0 or 1.
9. What is a Random Walk with Drift? Describe how it models time series data and give an example of its application in financial markets.
10. Critique the usefulness of the Signal in Noise model in various fields such as economics, engineering, and environmental science. How might the assumptions of this model limit its application?
11. Critically evaluate the effectiveness of each model (MA, AR, Random Walk with Drift, Signal in Noise) in handling different types of time series data.

Smoothing a Time Series

12. Explain the purpose of smoothing in time series analysis. What are the general goals of this technique?
13. Describe the moving average smoothing method. How does this method use weights to smooth data, and what are the effects of changing these weights?
14. Discuss how the moving average method helps in reducing the impact of random fluctuations in the data. What challenges might arise when using this method on time series with trends or seasonality?
15. Explain what kernel smoothing is and how it uses a Gaussian kernel to weigh data points. How does the choice of bandwidth affect the smoothing?

16. Detail the Lowess regression method. How does this method determine the weights for smoothing and how do these weights contribute to the robustness against outliers?
17. Explain the role of the parameter f in the `lowess()` function in R. How does changing the value of f affect the results of the Lowess smoothing?
18. Define smoothing splines and describe how they fit a spline function to the data. What does the regularization term in the loss function achieve?
19. Compare and contrast the advantages and potential drawbacks of using moving average, kernel smoothing, Lowess regression, and smoothing splines. When might one method be preferred over the others based on the characteristics of the time series data?

Time Series Regression

20. Define time series regression and explain how it differs from other types of time series analysis such as autoregressive models.
21. Describe the significance of including time as a variable in the regression models. What does this imply about the data and its relationship over time?
22. Discuss the use of lagged variables in time series regression. What are the benefits of including lagged terms?

Stationarity

23. Explain the difference between strict and weak stationarity. Why is weak stationarity more commonly used in statistical analysis of time series?
24. Elaborate on the impact of non-stationarity on the predictive performance of time series models. How does failing to account for stationarity potentially mislead forecasting?
25. Critique the assumption of constant variance in the definition of weak stationarity. How might changes in variance over time affect the validity of time series models?
26. Explain how the concept of weak stationarity might still inadequately describe the nature of certain financial time series. What alternative forms of stationarity might be considered?
27. Discuss how the mean, variance, and autocovariance function must behave for a time series to be considered weakly stationary.
28. Define autocovariance and autocorrelation. How are these metrics useful in analyzing the properties of a time series?
29. Analyze the implications of having a high autocorrelation at large lags for a given time series. What might this indicate about the underlying data generation process?
30. What does it indicate if the ACF values are outside the 95% confidence interval? How does this help in determining whether a time series is white noise?
31. Describe how you would assess the stationarity of a time series using graphical methods in R. What plots would you use and what features would you look for?

Dealing with Non-Stationarity

32. Discuss how the Box-Cox transformation generalizes other forms of transformations like logarithmic and square root transformations. What is the significance of the parameter λ in this transformation?
33. Explain the statistical reasoning behind using logarithmic transformations for time series data. What types of data characteristics make this transformation particularly effective?
34. Explain the process of detrending a time series. Why is it necessary, and how does it differ from differencing?
35. Describe the impact of detrending and differencing on the forecasting accuracy of a time series model. How might these preprocessing steps improve or impair the model's performance?
36. Provide a detailed explanation of the first and second differences of a time series. Under what circumstances might second differencing be necessary?
37. Explain how the autocorrelation function (ACF) can be used to verify the effectiveness of detrending and differencing interventions on a time series.

ARIMA Models

38. Define an ARIMA model and explain the components of its notation: ARIMA(p, d, q).
39. Describe a moving average model of order q , MA(q). How does it model the current value of the series?
40. Describe the structure of an autoregressive model of order p , AR(p). What does it mean for the model to have "memory" or "persistence"?
41. Discuss the role of the differencing operator ∇ in making a time series stationary. How does this relate to the integrated component of an ARIMA model?
42. Explain the purpose of the backshift or lag operator B in the context of ARIMA models. Provide an example of how it is used to define the differencing of a series.
43. Explain how the autoregressive operator $\phi(B)$ is used to form the equation of an AR(p) model.
44. Explain the significance of the moving average operator $\theta(B)$ in an MA(q) model.
45. Describe the combined model ARMA(p, q) and how it integrates features of both AR and MA models.
46. Explain how the properties of the ACF and PACF can help in selecting an appropriate ARIMA model for a time series. Provide examples of what the ACF and PACF might look like for different models.

Fitting an ARIMA Model

47. Detail how the orders of the AR and MA components (p and q) are identified using the ACF and PACF plots.
48. Explain the importance of model diagnostics in the ARIMA modeling process. What are some key diagnostic checks that should be performed?

49. Explain how information criteria such as AIC, AICc, and BIC are used to compare the fit of different ARIMA models. What does each criterion take into account?
50. Detail the process and importance of conducting model diagnostics after fitting an ARIMA model. What specific plots and statistics are typically used?
51. Discuss the implications of the Ljung-Box test results when diagnosing the fit of an ARIMA model. What does a significant result suggest about the residuals?

General Autoregressive Conditional Heteroscedasticity (GARCH) Models

52. Define an ARCH and a GARCH model and explain their importance in financial econometrics.
53. Discuss how a GARCH models can account for volatility clustering in financial time series.
54. Describe the basic structure of an ARCH(1) model and how it models the variance of a time series.
55. Explain how an AR(1) model with ARCH(1) error terms is constructed. Include a description of how each component contributes to modeling the time series.
56. Explain the extension from an ARCH model to a GARCH model. What additional features does a GARCH model incorporate?
57. Describe how to interpret the output of a fitted GARCH model, including parameter estimates and their significance.
58. Discuss the significance of the parameters α and β in a GARCH(1,1) model. What does each parameter represent, and how do they affect the model's behavior?

Chapter 13

Introduction to Neural Networks and Deep Learning

Sources and Further Reading

The material in this chapter is based on the following sources.

Gareth James, Daniel Witten, Trevor Hastie and Robert Tibshirani: *An Introduction to Statistical Learning with Applications in R*. 2nd edition, corrected printing, June 2023. (ISLR2)

<https://www.statlearning.com>

Chapter 10

While the James et al. book is otherwise very comprehensive, it only provides a single chapter on neural networks. The benefit here is that neural networks are discussed in context, as just one other regression or classification method. The downside is that it does not dive sufficiently deep into neural network architectures and fitting of neural network models that is at the heart of modern machine learning applications.

Kevin P. Murphy: *Probabilistic Machine Learning – An Introduction*. MIT Press 2022.

<https://probml.github.io/pml-book/book1.html>

Chapter 13, 14, 15

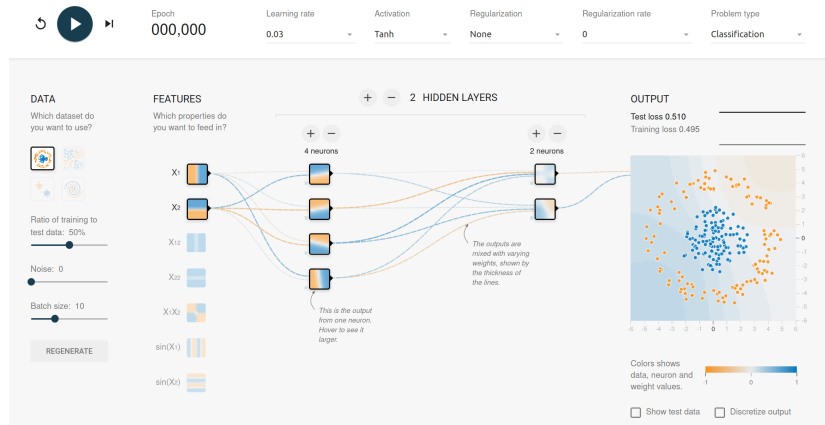


Figure 13.1: Tensorflow Playground

The book by Murphy is freely available under a Creative Commons license and provides three chapters on neural networks, one for structured data, one for images, and one for sequences. It provides significant depth on convolutional and recurrent network architectures, fitting the models, and problems the data analyst may encounter. However, it tends to focus on the mathematical background, rather than application or implementation.

Tensorflow Guides: <https://www.tensorflow.org/guide>

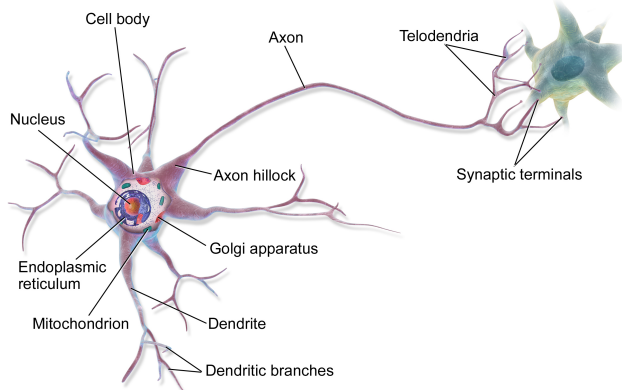
This section uses the Tensorflow programming framework for implementing neural network machine learning applications. The Tensorflow website has a multitude of introductory and advanced guides and tutorial that cover all aspects of machine learning with neural networks and are all very accessible to the beginner. While they focus heavily on implementation aspects, they provide significant coverage of the concepts as well.

Tensorflow Playground: <https://playground.tensorflow.org>

The Tensorflow Playground, shown in Figure 13.1, is a very visual and interactive introduction to how neural networks function. It allows playful exploration of a number of features with a small simulated neural network.

13.1 Introduction

Artificial Neural Networks ("ANN") are a type of non-linear statistical model for regression and classification. Their original motivation is the architecture of biological brains whose elementary unit is the *neuron*. Figure 13.2 shows an image of a biological



https://commons.wikimedia.org/wiki/File:Blausen_0657_MultipolarNeuron.png

Figure 13.2: Image of a biological neuron and its connections

neuron and its connections. Biological neurons in a brain are connected to many other neurons via *axons* that connect to other neurons at their *synapses*. Neurons receive electro-chemical inputs from other neurons through their axons. Once a certain threshold of input is reached, neurons themselves generate an electro-chemical potential that is transmitted to other neurons via their synapses. However, while this is the original motivation for ANNs, the architecture of modern ANNs is not modelled after biological brain architectures and ANNs are best understood as non-linear statistical models. From now on, the term "*neural network*" is used synonymously with "artificial neural network".

A basic neural network cell or unit is defined by a simple non-linear equation. *Inputs* x_i are weighted by *weights* w_i , then summed. A *bias* term b is added and the result is then transformed by a non-linear *activation function*.

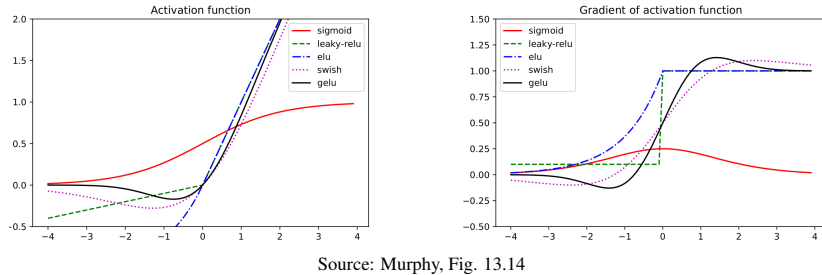
$$y = \psi(b + \sum_i w_i x_i) \quad (13.1)$$

It is important that the activation function is non-linear. Otherwise, even a complex network of such units would be nothing but a linear system and therefore equivalent in capabilities to a linear regression model. In other words, it is the non-linear activation functions that make neural networks more capable or more powerful than simple linear models. In addition to allowing a neural network to fit complex functional forms, the activation functions also serve to normalize, clip, or otherwise constrain the outputs of the neural unit and the entire network.

A great many activation functions have been proposed and investigated over the years. Table 13.1 shows frequently used ones and Figure 13.3 shows a selection of these functions and their gradient, that is, their first derivative. The most frequently-used

Sigmoid	$\sigma(z) = \frac{e^z}{1+e^z}$
Hyperbolic tangent	$\tanh(z) = \frac{\sinh(z)}{\cosh(z)} = \frac{e^z - e^{-z}}{e^z + e^{-z}} = 2\sigma(2z) - 1$
Softplus	$\sigma_+(z) = \log(1 + e^a)$
Rectified linear unit	$\text{ReLU}(z) = \max(a, 0)$
Leaky ReLU	$\text{LReLU}(z) = \max(z, 0) + \alpha \min(z, 0)$
Exponential linear unit	$\text{ELU}(z) = \max(z, 0) + \min(\alpha(e^z - 1), 0)$
Swish, Sigmoid linear unit	$\text{SiLU}(z) = z\sigma(z)$
Gaussian error linear unit	$\text{GeLU}(z) = z\Phi(z)$

Table 13.1: Selection of frequently-used activation functions



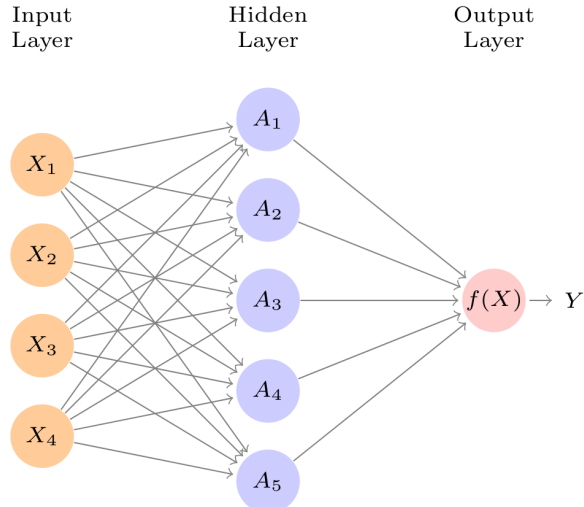
Source: Murphy, Fig. 13.14

Figure 13.3: Activation functions and their gradients

functions, and often the defaults in software implementations, are the ReLU, the tanh, and the sigmoid function. Note that a single neural unit with a sigmoid activation function is equivalent to a logistic regression model as seen in an earlier section.

The basic neural network units are typically arranged in *sequential layers*. The simplest form of a neural network is one with a single, fully-connected, hidden layer, shown in Figure 13.4. The network in Figure 13.4 has four inputs in its input layer, labelled $X_1 \cdots X_4$ and has a single cell or unit in its output layer. The layer of cells shown in blue and labelled $A_1 \cdots A_5$ are called “*hidden*” because they are neither input to the network, nor observable output. The layer is called “*fully-connected*” because each of the units in the layer is connected to all units of the previous layer, in this case the input layer. These characteristics make the network suitable to model a non-linear regression of one output on four inputs.

Recall the definition of a neural network unit in terms of its weights and biases in Equation 13.1. In the model in Figure 13.4, each connection from an input to a neural unit receives a weight w , and each neural unit adds a bias term b to the sum of its inputs. Counting the arrows and the units in Figure 13.4, the network has 25 weights (20 in the fully-connected hidden layer and 5 in the output layer) and 6 biases (5 in



Source: ISLR2 Figure 10.1

Figure 13.4: Neural network with a single fully-connected hidden layer

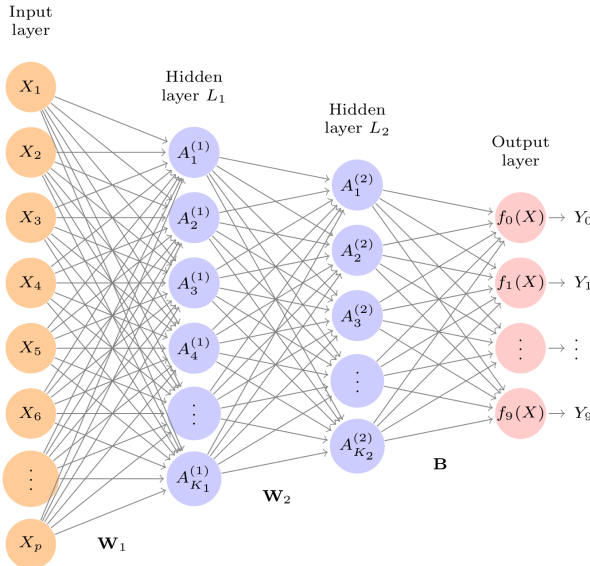
the fully-connected hidden layer and 1 in the output layer) for a total of 31 trainable or learnable parameters.

The basic network architecture of Figure 13.4 can be readily extended to multiple hidden layers and multiple output units in the output layer, as shown in Figure 13.5. This network has p inputs, K_1 units in the first hidden layer, K_2 units in the second hidden layer, and 10 units in the output layer. Again counting the connections and the units, this network has $p \times K_1 + K_1 \times K_2 + K_2 \times 10$ weights and $K_1 + K_2 + 10$ biases as trainable or learnable parameters.

An output layer with more than one output unit, such as the one in Figure 13.4 can be used for *multi-objective regression* analysis where more than one value is to be predicted jointly from the inputs. Another use of multiple outputs is *multi-class classification*, where each output represents the probability for a particular class. This is very similar to the multinomial logistic regression discussed in an earlier chapter. For multi-class classification in neural networks, the final outputs ("logits") are transformed to class membership probabilities using a *softmax* activation function:

$$\Pr(Y = m|X) = \frac{e^{Z_m}}{\sum_{l=0}^n e^{Z_l}}$$

where the Z_m are the logits, that is, the weighted sum of inputs plus the bias term, for class m of a total of n possible classes.



Source: ISLR2 Figure 10.4

Figure 13.5: Neural network with two fully-connected hidden layers and multiple outputs

13.2 Parameter Estimation

In order to learn (estimate) the parameters of a neural network, that is, to train the neural network, a *loss function* must be defined. This is the function that serves as the minimization objective and defines the difference between the outputs of the neural network and the target values, that is the correct or observed values. Typical loss functions for a regression analysis are the MSE (mean squared error), MAE (mean absolute error), Huber loss (a combination of MSE and MAE), or the MAPE (mean absolute percentage error). Typical loss functions for a classification analysis are the cross-entropy or KL-divergence after a softmax activation on the multiple output units.

As discussed above, the trainable or learnable parameters are the weights w and biases b of the units in the network. Together, they form the parameter vector that is defined as $\phi = (w, b)$.

13.2.1 Gradient Descent

Unlike the simple case of linear regression, there are no closed-form algebraic optimal solutions for the parameters available for general neural networks. Therefore, optimization is performed numerically using *stochastic gradient descent* (SGD). We first explain the simple non-stochastic gradient descent process.

The *gradient* is simply the vector of partial first derivatives of a function with multiple

inputs and is designated by the "nabla" symbol ∇ :

$$\nabla f(x_1, \dots, x_m) = \begin{pmatrix} \frac{\partial f}{\partial x_1} \\ \vdots \\ \frac{\partial f}{\partial x_m} \end{pmatrix}$$

For a simple function of one variable, like $f(x) = x^2$, the gradient is simply the first derivative:

$$\nabla f(x) = \frac{\partial f}{\partial x} = \frac{df}{dx} = 2x$$

As gradients are functions again, they can be evaluated for different inputs v , written as $\nabla f(x)|_v$. For example, the gradient of x^2 evaluated at $x = 2$ is $\nabla f(x)|_{x=2} = 4$ and the gradient of x^2 evaluated at 3 is $\nabla f(x)|_{x=3} = 6$.

Gradient descent is an iterative method to find the minimum of a function of multiple inputs:

1. Begin with random initial parameter values
2. Repeat the following two steps until convergence:
 - (a) Find the direction of steepest descent. This is the largest decrease in loss function value and is given by the gradient vector ∇L of partial derivatives.
 - (b) Move a step in that direction by adjusting the parameters. The step size is determined by the *learning rate*)

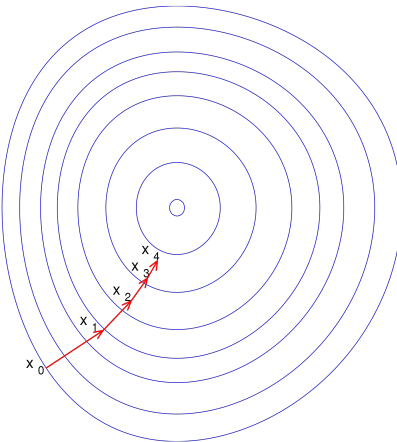
This procedure can be summarized as follows. Consider the loss L at a certain input X as a function of parameter values θ . Then, at each step t , update the parameters θ using learning rate γ until the parameter values do not change anymore (in practice: until the changes are smaller than a certain threshold):

$$\theta_{t+1} = \theta_t - \gamma \nabla L(\theta)|_{\theta_t, X} \quad (13.2)$$

Recall that the vertical bar notation in the final term means "evaluated at", that is, the gradient of $L(\theta)$ is evaluated at input values X and current parameter values θ_t .

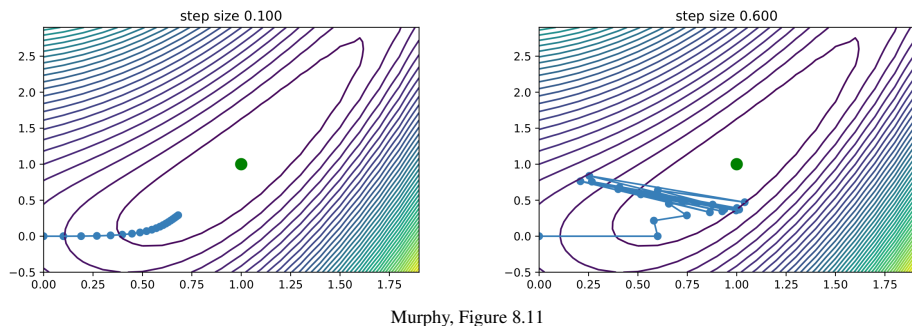
Figure 13.6 shows an image of such a process. Beginning with an initial solution, the process of gradient descent will eventually arrive at the optimal solution.

Numerical optimization via gradient descent is prone to a number of potential problems, among them slow convergence, no convergence (oscillations) and premature



https://commons.wikimedia.org/wiki/File:Gradient_descent.svg

Figure 13.6: Illustration of gradient descent



Murphy, Figure 8.11

Figure 13.7: Slow convergence and no convergence in gradient descent

convergence to a local optimum rather than the global optimum. The first two are illustrated in Figure 13.7.

In the left panel of Figure 13.7, the gradient of a function of two parameters (vertical and horizontal axis) is very "shallow", that is, the function changes values slowly with the parameters and the gradient is very small. When the step size λ is too small, it will take very many steps to reach the optimum.

One might imagine that the solution would be to simply increase the step size. However, the right panel of Figure 13.7 shows a lack of convergence that may be due to a large step size. Because the gradient is so shallow, the gradient is similar in multiple directions. Once a gradient descent step takes the parameter vector into a steeper region of the gradient, the next step is likely to "overshoot" the correction, leading to back-and-forth oscillations shown in the right panel of Figure 13.7. The gradient descent

process will not converge to the optimum.

Finally, there may be functions not only with a global optimum but with multiple local optima. In those cases, it is possible that the gradient descent process will converge to a local optimum, that is, it will show premature convergence. Simply increasing the step size to get out of the local optimum is not a solution, as it is often unclear whether the optimum that is found is local or global and a larger step size can lead to the earlier oscillation problem.

13.2.2 Stochastic Gradient Descent

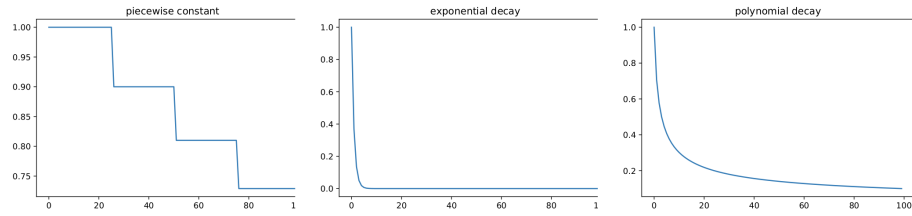
Note that in Equation 13.2 the gradient is evaluated at the current parameters θ_t and the inputs to the neural network X . In practice, this means that the gradient should be computed as the mean gradient over all training observations. With large training sets, this is computationally expensive and wasteful, as it is likely that a small sample of the input can already point the gradient descent process in the right direction.

This is the motivation behind *stochastic gradient descent*. Instead of computing the gradient and averaging over the entire input training set, gradient updates of the form in Equation 13.2 are done for "*minibatches*" that are randomly sampled from the training data set. The gradient is evaluated for each observation of the minibatch and then averaged over the minibatch. A typical minibatch size is anywhere between 16 and 256 observations. Minibatches should be randomly drawn and independent of each other.

The processing of all minibatches in the training data set is called an "*epoch*". Typically, a neural network is trained for multiple epochs, until the parameters converge to their optimal values. To avoid repetition of inputs, the training data should be randomly shuffled between epochs.

The decision as to how many training epochs should be used can be made in different ways. The simplest is to use a fixed, large number of episodes with the hope that training will have converged to an optimum at the end of the procedure. However, this simple method may be computationally wasteful if training converges rapidly, and it may also lead to overfitting. *Early stopping* is typically used to prevent this. Here, the data set is split into training, validation, and test samples. The training samples are used for training. After each epoch, the validation set is used to assess the prediction error. When the prediction errors are stable, that is, the changes in validation errors after a number of successive epochs are smaller than a given threshold, training is stopped. The prediction error of the final model is then evaluated using the test data set. Early stopping thus serves as a *regularization method* that prevents overfitting, and also saves computational resources, but at the expense of reducing the size of the training and/or the test data set (because a separate validation sample is required).

Because each gradient update step (Equation 13.2) is evaluated at a different input X , the gradient can be different from update step to update step, even if the parameters are largely unchanged. Hence, especially with small minibatch sizes, the gradient can be highly unstable, and the gradient descent proceeds in more or less random directions,



Source: Murphy Figure 8.18

Figure 13.8: Adaptive learning rates

further adding to the potential convergence problems note above.

13.2.3 Parameter Updates

In order to tackle the convergence issues in SGD, a number of solutions have been proposed. The simplest of these is to use an *adaptive learning rate*, where the learning rate is large in early training and then decreases. The idea is to accelerate early training and prevent oscillations or "overshooting" the optimum later in training. Figure 13.8 shows three examples of adaptive learning rates. The left panel shows a piece-wise constant learning rate that is periodically decreased, the center panel shows an exponential decay of the learning curve, and the right panel shows a polynomial decay in the learning curve. Which approach is best depends on the specifics of the problem. It is not unusual for a business analyst to use small subsamples of the training data to try different methods and evaluate their convergence behaviour before training on the full training data set.

A more sophisticated way is the *AdaGrad* method ("adaptive gradients") that was originally developed for sparse gradient vectors, that is, gradients with many 0 components. This method scales the learning rate inversely to the square root of the sum of squared gradients of all previous steps (Equations 13.3 and 13.4). The intuition is that large gradients should lead to small updates, and vice versa. The overall learning rate can also be adjusted but is typically left constant ($\lambda_t = \lambda$). The term ϵ in Equation 13.3 is a very small value to prevent division by zero in case all the s_t are 0 (or close to it). Note that because the sum of previous squared gradients is monotonically increasing, the learning rate monotonically decreases — it can never increase.

$$\theta_{t+1} = \theta_t - \lambda_t \frac{1}{\sqrt{s_t + \epsilon}} \nabla L(\theta)|_{\theta_t, X} \quad (13.3)$$

$$s_t = \sum_{\tau=1}^t (\nabla L(\theta)|_{\theta_\tau, X})^2 \quad \text{Sum of all prior squared gradients} \quad (13.4)$$

The *RMSProp* ("Root Mean Square Propagation") modifies AdaGrad to overcome its monotonically decreasing learning rate and to prevent a learning reduction that is too

quick. Instead of accumulating all past squared gradients, RMSProp uses an exponential moving average that emphasizes recent gradients and makes it more robust to large changes in gradients. Note how the square of previous gradients is propagated by the term s_t on the right hand side of Equation 13.5. Equation 13.5 is used in place of Equation 13.4 in the update equation (Equation 13.3).

$$s_{t+1} = \beta s_t + (1 - \beta) (\nabla L(\theta)|_{\theta_t, X})^2 \quad (13.5)$$

The *AdaDelta* method is an extension of RMSProp that seeks to reduce its dependence on a global learning rate λ . Instead of using λ , AdaDelta uses the ratio of the root mean square of previous parameter updates to the root mean square of the current gradient. This method adapts learning rates based on parameter update history, represented by $\Delta\theta_t$ (Equation 13.6), providing a more stable and responsive adjustment mechanism. The term s_t in Equation 13.6 is that of Equation 13.5.

$$\begin{aligned} \theta_{t+1} &= \theta_t + \Delta\theta_t \\ \Delta\theta_t &= -\lambda_t \frac{\sqrt{\delta_{t-1} + \epsilon}}{\sqrt{s_t + \epsilon}} \nabla L(\theta)|_{\theta_t, X} \\ \delta_t &= \beta\delta_{t-1} + (1 - \beta)(\Delta\theta_t)^2 \end{aligned} \quad (13.6)$$

While Adagrad, RMSProp, and AdaDelta method all modify the parameter update step sizes to allow a gradual reduction in learning rate in order to tackle problems of slow or premature convergence, another way to look at these methods is as variance reduction methods of the parameter or parameter updates between minibatches. In other words, the variability of the parameter updates (and therefore, the variability of the parameters) from training batch to training batch is reduced. This then tackles the problems of oscillations or rapid changes in the direction of parameter updates illustrated in the right panel of Figure 13.7.

Momentum methods for SGD are another way to address rapid changes in parameter updates. They are based on the intuition of physical momentum, that is, to keep going in the general direction of previous updates and any changes have only small effects on this direction, that is, to avoid "sharp turns" in the direction of the parameter changes. This is also intended to avoid the oscillations that can prevent convergence. The standard momentum method is defined as:

$m_{t+1} = \beta m_t - \nabla L(\theta) _{\theta_t, X}$	Momentum
$\theta_{t+1} = \theta_t - \lambda m_{t+1}$	Parameter update

Typical values for β are ≈ 0.9 and good values for β must again be found by experimenting with subsamples and observing convergence behaviour.

The *Nesterov momentum*, a variant of the standard momentum method, incorporates a look-ahead step to the parameter updates, making the convergence faster and more responsive to the loss surface. Note how the gradient in Equation 13.7 is evaluated at $\theta_t + \beta m_t$ — it is evaluated not at the current parameter values but at those after the next update step, as defined in Equation 13.8.

$$m_{t+1} = \beta m_t - \lambda_t \nabla L(\theta)|_{\theta_t + \beta m_t, X} \quad \text{Nesterov Momentum} \quad (13.7)$$

$$\theta_{t+1} = \theta_t + m_{t+1} \quad \text{Parameter update} \quad (13.8)$$

Finally, the *AdaM* technique (Adaptive Moment Estimation) combines ideas from both AdaGrad and RMSProp, adjusting learning rates based on both the gradient and the squared gradient:

$$m_t = \beta_1 m_{t-1} + (1 - \beta_1) \nabla L(\theta)|_{\theta_t, X} \quad (13.9)$$

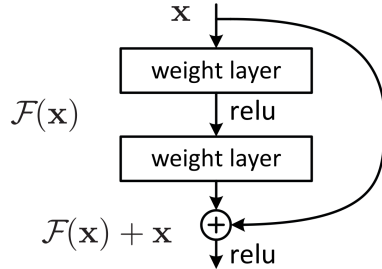
$$s_t = \beta_2 s_{t-1} + (1 - \beta_2) (\nabla L(\theta)|_{\theta_t, X})^2 \quad (13.10)$$

$$\theta_{t+1} = \theta_t - \lambda_t \frac{1}{\sqrt{s_t} + \epsilon} m_t \quad (13.11)$$

13.2.4 Gradient Problems

The issue of *vanishing gradients* arises when gradients of the network's weights become very small, effectively preventing weights from changing their values despite large step sizes. As gradients of the loss function are propagated backwards through the network to update weights, the gradient values diminish with each layer as small derivatives are multiplied with other small derivatives. The gradients can become so small that they have virtually no effect in updating some layers, especially those layers close to the input. This problem is increasingly likely to occur the more layers the network contains. Activation functions like the sigmoid or the tanh function contribute to this problem because they asymptotically restrict values between 0 and 1. The gradients at the left and right extremes are very small, the activation functions there are shallow as seen in Figure 13.3. There are a number of solutions to tackle this problem:

- Using non-saturating activation functions, that is, functions that do not asymptotically approach a fixed value, like ReLU (Rectified Linear Unit) or its variants (e.g., Leaky ReLU, ELU) is effective. The ReLU does not asymptotically approach a constant and its gradient therefore does not approach zero for large inputs.
- Architectures such as *Residual Networks* (ResNets) help mitigate the vanishing gradient problem by incorporating "skip connections" that allow gradients to flow through an alternate shortcut path, bypassing several layers at a time. Figure 13.9 shows an example of a ResNet architecture, where the \oplus sign indicates addition of vectors.



Source: Murphy, Figure 13.15

Figure 13.9: ResNet architecture

- *Batch normalization* standardizes the inputs to a layer within a network. This helps maintain a mean output close to zero where the gradients of sigmoid and tanh functions are largest, preventing the gradients from vanishing too quickly during training.
- Suitable initialization of weights can help in preventing the vanishing gradient issue at the start of training. Initial values for neural network parameters θ_0 are typically drawn randomly from a normal distribution:

$$\theta_0 \sim N(0, \sigma^2)$$

A number of different ways of setting the variance σ^2 of this distribution are used in practice. *Glorot initialization* (also called *Xavier initialization*) sets the variance as a function of the number of incoming connections n_{in} and the number of outgoing connections n_{out} is from each unit:

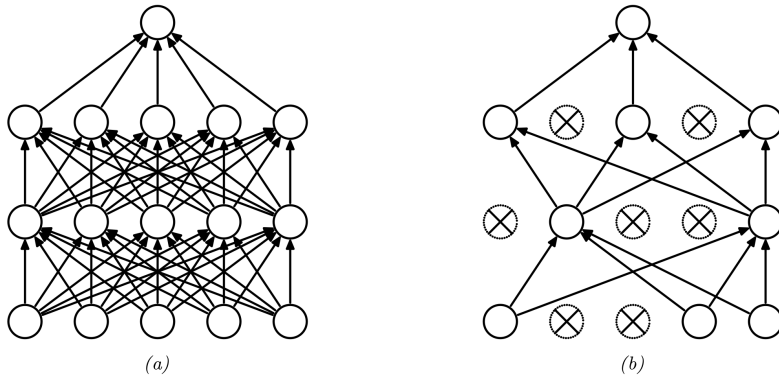
$$\sigma^2 = \frac{2}{n_{\text{in}} + n_{\text{out}}}$$

In contrast, LeCun initialization focuses only on the number of inputs to each unit:

$$\sigma^2 = \frac{1}{n_{\text{in}}}$$

and He initialization doubles the variance of the LeCun initialization:

$$\sigma^2 = \frac{2}{n_{\text{in}}}$$



Source: Murphy, Figure 1.318

Figure 13.10: Example of dropout regularization in neural networks

Which of these results in the best learning performance is problem-specific and depends on the type of neural network model, the loss function, and the data set. It is typically explored experimentally with small subsamples of the training data for which optimization progress is observed.

The opposite problem of vanishing gradients is that of *exploding gradients*. Gradients can grow exponentially across layers as they are propagated through the network due to the cumulative multiplication of large derivatives with other large derivatives. This often results in the model parameters being updated in ways that cause the learning process to diverge, leading to learning instability and oscillations of parameter estimates. *Gradient clipping* is the most direct method to combat exploding gradients. Gradient clipping involves setting a threshold value, and if the gradients exceed this value, they are simply limited or cut to that value.

13.2.5 Regularization with Dropout

In addition to early-stopping as a regularization method, using "*dropout*" has been shown to be effective in preventing overfitting a model. Dropout randomly, for each observation in a minibatch, removes a fraction of units in a layer, or, equivalently, randomly sets the output of a fraction of units to 0. The dropout effect mimics training a large ensemble of neural networks, each with a slightly different architecture, and each network trained on different subsets (minibatches) of data. Dropout is done for training only, not when evaluating the performance for the validation or test sample.

Figure 13.10 shows an example. In the right panel, the crossed-out units of the model in the right panel are dropped, that is, their output is set to 0. The intuition is that it prevents the model to use particular units or connections to represent specific observations in the training data. Dropout rates are commonly set to ≈ 0.25 of all units in a layer but is not uncommon to see rates as high as 0.50.

13.3 Software Frameworks for Neural Network Models

The landscape of neural network software frameworks has expanded significantly in recent years, driven by an increasing demand for more sophisticated machine learning solutions. These frameworks are designed to facilitate the design, training, and deployment of neural networks with high efficiency and flexibility. The choice of a neural network software framework depends largely on the specific needs of the project, including the ease-of-use, flexibility, and performance requirements, and the specific type of neural network being implemented. Each of the following frameworks offers unique strengths that cater to different aspects of neural network development and deployment. In the end, the choice of framework often comes down to developer familiarity and preference, and the need to fit into a larger software system and development team.

Tensorflow and Keras



TensorFlow, developed by the Google Brain team and released as version 1.0 in 2018, is an open-source framework designed for high-performance numerical computation, in particular for neural network applications. TensorFlow supports both CPU and GPU computing, and it can be scaled from single machines to large clusters, making it suitable for a wide range of applications, from developing simple models on a desktop to deploying complex systems in production environments. Tensorflow provides automatic differentiation and computation of gradients, supports distributed computing across clusters, and parallel computing on multiple GPUs per compute node. Tensorflow provides a wide range of loss functions, optimizers, activation functions and neural network architectures.



Keras is a high-level neural networks API, written in Python and originally developed for running on top of TensorFlow, Theano, or Microsoft Cognitive Toolkit (CNTK). Later releases of Keras focused on TensorFlow only. Keras aims to enable fast experimentation with deep neural networks, focusing on being user-friendly, modular, and extensible. It provides simple and consistent high-level APIs (application programming interfaces) that make it easy to create deep learning models without getting bogged down in the details of the underlying algorithms. Keras handles the specifics of creating complex network layers, loss calculations, and optimization steps, allowing developers to focus on building the core parts of their models. It provides a wide range of neural network layer types to support different architectures and offers simplified data management for training. Keras is officially integrated into TensorFlow as the `tf.keras` package, making it the default high-level framework for building and training machine learning models in TensorFlow.

PyTorch



prototyping.

Developed by Facebook's AI Research lab, PyTorch is known for its simplicity, ease of use, and dynamic computational graph that allows for mutable graph executions. This feature is particularly useful in research and development environments where iterative and explorative approaches are common. PyTorch also provides strong support for GPU acceleration and has gained a lot of popularity in the academic and research community for its ease in

Theano / PyTensor



Theano was originally at the University of Montreal. Although development has stopped, the project lives on as PyTensor and remains influential in the academic world. Theano or PyTensor are Python libraries that allows for defining, optimizing, and evaluating mathematical expressions involving multi-dimensional arrays efficiently. It is particularly well-known for its performance and precision in computations, which is why it was heavily used in academically oriented projects.

Microsoft Cognitive Toolkit (CNTK)



Also known as CNTK, this deep learning framework from Microsoft provides an efficient environment for training deep learning models at scale, across multiple GPUs and machines. CNTK supports both declarative and imperative programming languages and is known for its performance in handling large datasets.

13.4 Linear Regression using Tensorflow and Keras

Complete implementations for this and the other examples in this chapter are available on the following GitHub repo: <https://github.com/jevermann/busi4720-ml>

The project can be cloned from this URL: <https://github.com/jevermann/busi4720-ml.git>

As noted in the previous section, Keras makes the construction of a neural network model very easy, as the Python example in this section shows. This first example shows a linear regression in Keras. In particular, no non-linear activation functions are used.

First, the required packages are imported. The `pandas` package is required to read the training data, and the `tensorflow` package includes Keras as well:

```
import pandas as pd
import numpy as np
import tensorflow as tf
from tensorflow.keras import layers
```

Next, the training data set is read from a CSV file. This example uses the Boston housing price dataset that from the R package `ISLR2` was used previously in regression analysis examples:

```
# Use the Boston housing data set
boston_data = pd.read_csv("https://evermann.ca/busi4720/boston.csv")
```

The example uses all columns as input features, except the `medv` column, the median value of house prices in a suburb, which is the prediction target or the true label for the observation:

```
boston_features = boston_data.copy()
boston_labels = boston_features.pop('medv')
```

Next, the neural network model is defined. It is a model that contains two sequential layers, both fully-connected ("dense" in the Tensorflow/Keras terminology). Keras provides the `Sequential` model type for defining this, which accepts a list of layer definitions, here the `Dense` layer types:

```
boston_model = tf.keras.Sequential([
    tf.keras.layers.Dense(64, activation=None),
    tf.keras.layers.Dense(1, activation=None)
])
```

Keras provides many different model types and layer types, other than sequential layers to allow the analyst to define more complex neural network architectures. Consult the Keras documentation at <https://keras.io/api/models/> and <https://keras.io/api/layers/> for an overview and details.

Note that the input layer and its dimensions are not explicitly described in this example. When training this model, Keras will automatically adapt the first dense layer to the actual input data provided. Later examples will require specification of the input layers when the size of the input data cannot be determined automatically, or when layers cannot be dynamically scaled or adjusted.

The second dense layer serves as the output layer with a single output unit. Note that no activation function is specified for either layer. In other words, this example is a linear regression.

Each of the k input features is passed to each of the 64 units in the first layer. From Equation 13.1, this model will have $k \times 64 + 64$ weights and $64 + 1 = 65$ biases as trainable parameters. With $k = 12$ feature columns in `boston_features`, this model will have $832 + 65 = 897$ trainable parameters.

Model compilation using the `compile()` function in Keras is used to specify the loss function and the optimizer to use. As a regression analysis, this example uses the MSE as loss function, and the Adam optimizer for SGD.

```
boston_model.compile(  
    loss = tf.keras.losses.MeanSquaredError(),  
    optimizer = tf.keras.optimizers.Adam())
```

With model compilation complete, the model can be trained, that is, fitted to the data using the `fit()` function. This example trains the model for 25 epochs. When no batch size is specified, as in this example, Keras by default chooses 32 as batch size.

```
boston_model.fit(boston_features, boston_labels, epochs=25)  
boston_model.summary()
```

The `summary()` function does not show a summary of training results but a summary of the model architecture. It is only once the `fit()` function has been called that Keras knows the size of the input layer, that is, the number of feature columns, and therefore the exact architecture of the model. The output of the `summary()` function is reproduced below. The output of each layer is two-dimensional, with the first dimension (the "rows") being the batch size, and the second dimension (the columns) being the outputs of the neural units at that level. The word `None` in the output shapes of each layer indicate that the minibatch size is flexible. The model summary also confirms the number of trainable parameters, as explained earlier.

```
>>> boston_model.summary()  
Model: "sequential"  
  
-----  
Layer (type)           Output Shape        Param #  
=====-----  
dense (Dense)          (None, 64)           832  
dense_1 (Dense)         (None, 1)            65  
=====-----  
Total params: 897 (3.50 KB)  
Trainable params: 897 (3.50 KB)  
Non-trainable params: 0 (0.00 Byte)
```

Verifying the number of parameters and understanding what they do is important to understand and verify the correctness of the implemented model!

13.5 Non-Linear Regression using Tensorflow and Keras

The example in this section illustrates the use of non-linear activation functions, feature preprocessing (normalization), model fitting with a validation data set, additional training and validation metrics, and final visualization of model training. It provides a more realistic example of the use of Tensorflow and Keras.

For normalization of numerical features, Keras provides a Normalization pre-processing layer that can be included in the model:

```
norm_layer = tf.keras.layers.Normalization()
```

This layer scales the input features to have a mean of 0 and a standard deviation of 1. To do this, the layer needs to "take a look at" the training data to determine the column means that must be subtracted from the data, and the column standard deviations that it must divide the data by. This is done by calling the `adapt()` function of the normalization layer on the training data, first converted to a numpy array.

```
norm_layer.adapt(boston_features.to_numpy())
```

This example also uses a sequential model. The first layer is the normalization layer that has been adapted to the training data. The remaining hidden and output layers are similar to the earlier example, but this example uses a ReLU activation function for the hidden layer and leaves the output layer without an activation.

```
norm_boston_model = tf.keras.Sequential([
    norm_layer,
    tf.keras.layers.Dense(64, activation='relu'),
    tf.keras.layers.Dense(1, activation=None)
])
```

The model summary output below shows the normalization layer as part of the model. Note that a summary can be produced in this case prior to model fitting, because the normalization layer already "knows" about the shape of the data through its adaptation.

```
>>> norm_boston_model.summary()
Model: "sequential_1"
=====
Layer (type)          Output Shape         Param #
=====
normalization (Normalizati (None, 12)           25
dense_2 (Dense)        (None, 64)            832
dense_3 (Dense)        (None, 1)              65
=====
Total params: 922 (3.61 KB)
Trainable params: 897 (3.50 KB)
Non-trainable params: 25 (104.00 Byte)
```

The number of trainable parameters is the same as in the previous model. However, the normalization layer contains 25 parameters that are not trainable: the 12 columns means, the 12 column standard deviations, and one parameter for the number of observations used in `adapt()`.

Verifying the number of parameters and understanding what they do is important to understand and verify the correctness of the implemented model!

The loss and optimizer functions are set as in the previous example. However, the following code block also illustrates the use of `metrics` specified in the `compile()` function. These are additional metrics that Keras will compute during training and testing. They are accessible from the training history object that is returned by the `fit()` function.

```
# Define loss and specify optimizer
norm_boston_model.compile(
    loss = tf.keras.losses.MeanSquaredError(),
    optimizer = tf.keras.optimizers.Adam(),
    metrics = ['mse', 'mae'])
```

The `fit()` function returns a history of the loss function and any additional metrics specified in `compile()`. The following Python code block also illustrates specifying a different minibatch size (20, the default is 32), and a training/validation split for the sample. After every epoch, Keras will evaluate the trained model on the validation sample.

```
# Fit model to data
train_hist = norm_boston_model.fit(
    boston_features,
    boston_labels,
    batch_size=20,
    epochs=50,
    validation_split=0.33)
```

The final output of this example looks like the following, indicating the loss function values and the additional metric values for both training and validation data sets for each epoch:

```
Epoch 1/50
1/17 [>.....] - ETA: 8s - loss: 784.9610 -
mse: 784.961017/17 [=====] - 1s 19ms/step -
loss: 693.6062 - mse: 693.6062 - mae: 24.9111 - val_loss: 366.6743 -
val_mse: 366.6743 - val_mae: 17.3806
...
Epoch 50/50
1/17 [>.....] - ETA: 0s - loss: 8.3212 -
mse: 8.3212 - m17/17 [=====] - 0s 4ms/step-
loss: 13.6148 - mse: 13.6148 - mae: 2.8122 - val_loss: 200.7858 -
val_mse: 200.7858 - val_mae: 11.5683
```

The metrics can also be plotted. The following Python code example uses the Plotly Express package and produces the graph shown in Figure 13.11.

```
import plotly.express as px
import numpy as np
# Transform the training history into a suitable data frame
hist = pd.DataFrame({
    'training': train_hist.history['mse'],
    'validation': train_hist.history['val_mse']})
hist['epoch'] = np.arange(hist.shape[0])
hist = pd.melt(hist, id_vars='epoch',
               value_vars=['training', 'validation'])
# Plot training history
fig = px.line(hist, x='epoch', y='value', color='variable')
fig.show()
```

Hands-On Exercise

- Modify the above code to include different activation functions, e.g. "tanh", "sigmoid", or "relu". Comment on the learning progress and loss function values.
- Modify the above code to change the number of units in the dense layer. Comment on the learning progress and loss function values.
- Modify the architecture to add one or more dense layers with different numbers of units. Comment on the learning progress and loss function values.

13.6 Classification using Tensorflow and Keras

The classification example in this section uses the `wage` dataset from the ISLR2 library for R. The dataset has been adapted to include a column `wagequart`, the quartile of

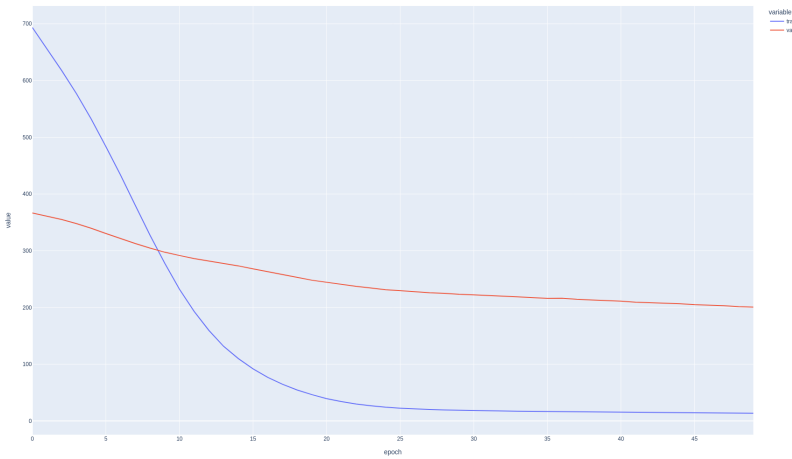


Figure 13.11: Regression example training and validation loss (MSE)

the wage to be predicted. Many features are categorical, encoded as strings, and need to be converted to numeric inputs. This example illustrates additional techniques beyond what the regression examples have demonstrated. These include categorical predictors and their pre-processing, as well as the use of TensorBoard for visualization of the model and the learning results.

First, read the data set and separate features and target labels, as in the regression example above:

```
# Read data and separate features from target labels
wage_data = pd.read_csv("https://evermann.ca/wage.csv")

wage_features = wage_data.copy()
wage_labels = wage_features.pop('wagequart') - 1
```

This example encodes each categorical feature using *one-hot encoding*. Similar to contrasts in traditional regression models, one-hot-encoding uses binary dummy variables to express the different categories. Unlike contrasts in traditional regression models, which use $k - 1$ dummy variables to encode k categories, one-hot encoding uses k dummy variables to encode k categories. There is no "base" category encoded by all dummy variables being zero, as in contrasts in traditional regression. Instead, every category is represented by its own dummy variable, and exactly one of the dummy variables is one.

Because there are many categorical variables in the data set, the transformation to one-hot is done using a `for` loop in the Python code block below. This example introduces the Keras `Input` object, which is used to specify the shape and type of input for

each categorical feature. The Keras StringLookup layer takes as input a character string denoting the category and turns character strings into one-hot dummy variables (or, alternatively) into integer category numbers. The layer must be adapted, that is, the data set is used to determine the number of different categories and therefore the number of binary dummy variables for that layer. After adaptation, the lookup layer is applied to the input object to create the one-hot encoded feature. The Keras input object is then added to a dictionary of input objects, and the pre-processed, one-hot encoded feature is added to a list pre-processed.

```
# Keep track of inputs and preprocessed inputs
inputs = {}
preproc_inputs = []

# Loop over all categorical features
for cat_feature in ['marital', 'race', 'education', \
                     'jobclass', 'health', 'health_ins']:

    # An Input variable is a placeholder that
    # accepts data input when training or predicting
    input = tf.keras.Input(shape=(1,),
                           name=cat_feature, dtype=tf.string)

    # This StringLookup layer accepts a string and
    # outputs its category as a one-hot vector (or,
    # alternatively as an integer)
    lookup = tf.keras.layers.StringLookup(
        name=cat_feature+"_lookup", output_mode="one_hot")

    # Adapt it to the different strings in the data
    lookup.adapt(wage_features[cat_feature])

    # And tie the input to this layer
    onehot = lookup(input)

    # Add the input feature to the list of inputs
    inputs[cat_feature] = input
    # Append the preprocessed feature to list of preprocessed inputs
    preproc_inputs.append(onehot)
```

As in the earlier example, numerical input features (here, only the feature `age`) are normalized using a normalization layer. The layer is adapted to the data set (the column "age" of the Pandas dataframe). Next, the normalized age feature is formed by applying the normalization layer to the age input object. The "age" input object feature is then added to the dictionary of input objects, and the normalized feature is appended to the list of pre-processed inputs.

```

age_input = tf.keras.Input(shape=(1,), name="age", dtype="float32")
norm_layer = tf.keras.layers.Normalization(name="age_norm")
norm_layer.adapt(wage_features["age"])
age_norm = norm_layer(age_input)

inputs["age"] = age_input
preproc_inputs.append(age_norm)

```

The integer feature `year` will also be treated as categorical, rather than numerical. For this, Keras provides an `IntegerLookup` layer, which behaves analogous to the `StringLookupLayer` above but for integer inputs instead of character string inputs.

```

# Define the input placeholder
year_input = tf.keras.Input(shape=(1,), name="year", dtype="int32")

# Define and adapt an IntegerLookup layer for the one-hot encoding
lookup = tf.keras.layers.IntegerLookup(name="year_lookup",
                                       output_mode="one_hot")
lookup.adapt(wage_features["year"])
year_onehot = lookup(year_input)

# Add the input and preprocessed input to the collections
inputs["year"] = year_input
preproc_inputs.append(year_onehot)

```

The next step concatenates all preprocessed input features into a single vector using a Keras `Concatenate` layer:

```

preprocessed_inputs = \
    tf.keras.layers.Concatenate(name="concat") (preproc_inputs)

```

The next step builds a first Keras model that is responsible only for the preprocessing of inputs. Instead of using a `Sequential` model type construction, a Keras *Model* object is created directly by supplying its inputs (`inputs`) and outputs (the `preprocessed_inputs`) as well as a name for the model. Note that in defining the preprocessing in the above code blocks, the outputs can be traced back to the preprocessing layers and then to the corresponding inputs; Keras assembles a model using this information.

```

preproc_model = tf.keras.Model(inputs,
                               preprocessed_inputs,
                               name="preproc")
preproc_model.summary()

```

The summary for this model is printed below. Note the output shape of the input layers are one element each (that is, the character string expressing the category, or the

integer for the year, or the floating point value for the age), whereas the output of the corresponding string lookup layers are one-hot-encoded and have as many elements as there are categories for that feature. Also note last column that shows the connections between a layer and its input. Finally, except for the normalization layer for age, no other layer has any parameters. The age normalization layer has three parameters, corresponding to the mean, standard deviation, and number of observations of the inputs it has received during adaptation. These are non-trainable parameters and their values are determined from the data when the layer was adapted.

```
>>> preproc_model.summary()
Model: "preproc"

Layer (type)          Output Shape Param Connected to
=====
maritl (InputLayer)  [(None, 1)] 0      []
race (InputLayer)   [(None, 1)] 0      []
education (InputLayer) [(None, 1)] 0      []
jobclass (InputLayer) [(None, 1)] 0      []
health (InputLayer) [(None, 1)] 0      []
health_ins (InputLayer) [(None, 1)] 0      []
age (InputLayer)   [(None, 1)] 0      []
year (InputLayer)  [(None, 1)] 0      []
maritl_lookup (StringLo (None, 6) 0      ['maritl[0][0]']
race_lookup (StringLook (None, 5) 0      ['race[0][0]']
education_lookup (Strin (None, 6) 0      ['education[0][0]']
jobclass_lookup (String (None, 3) 0      ['jobclass[0][0]']
health_lookup (StringLo (None, 3) 0      ['health[0][0]']
health_ins_lookup (Stri (None, 3) 0      ['health_ins[0][0]']
age_norm (Normalization (None, 1) 3      ['age[0][0]']
year_lookup (IntegerLoo (None, 8) 0      ['year[0][0]']
concat (Concatenate)    (None, 35) 0      ['maritl_lookup[0][0]',
                                             'race_lookup[0][0]',
                                             'education_lookup[0][0]',
                                             'jobclass_lookup[0][0]',
                                             'health_lookup[0][0]',
                                             'health_ins_lookup[0][0]',
                                             'age_norm[0][0]',
                                             'year_lookup[0][0]']

=====
Total params: 3 (16.00 Byte)
Trainable params: 0 (0.00 Byte)
Non-trainable params: 3 (16.00 Byte)
```

Verifying the output shapes of each layer is important to understand and verify the correctness of the implemented model!

With the preprocessing complete, the actual classification model can be built. The Python code block below defines this as a three-layer sequential model with four output units, corresponding to the four classes to be predicted. The activation on the output layer is a softmax so that the model outputs class-membership probabilities, rather than

logits (as an equivalent, Keras provides an explicit `Softmax` layer that could be used). Instead of providing layers as a list to the Keras `Sequential` model constructor as in the earlier example, this example uses the `add()` function to add layers to the model.

```
class_model = tf.keras.Sequential(name="classification")
class_model.add(tf.keras.layers.Dense(64, activation="relu"))
class_model.add(tf.keras.layers.Dropout(0.25))
class_model.add(tf.keras.layers.Dense(32, activation="relu"))
class_model.add(tf.keras.layers.Dropout(0.25))
class_model.add(tf.keras.layers.Dense(4, activation="softmax"))
# Alternatively:
# class_model.add(tf.keras.layers.Dense(4, activation=None))
# class_model.add(tf.keras.layers.Softmax())
```

To connect the model for feature preprocessing and the classification model, it is important to recall that the output of the preprocessing model is the input to the classification model. That is, the output of the preprocessing model when applied to the `inputs`, is the input to the `class_model`. In other words, the classification result is the output of applying the class model to the output of the preprocessing model when applied to the input objects, as shown in the following Python code block:

```
class_results = class_model(preproc_model(inputs))
class_model.summary()
```

The output of the `summary()` function is shown below. The reader should verify the number of trainable parameters and understand what these parameters represent in the model.

Model: "classification"			
Layer (type)	Output Shape	Param #	
dense_4 (Dense)	(None, 64)	2304	
dropout (Dropout)	(None, 64)	0	
dense_5 (Dense)	(None, 32)	2080	
dropout_1 (Dropout)	(None, 32)	0	
dense_6 (Dense)	(None, 4)	132	

Total params: 4516 (17.64 KB)
Trainable params: 4516 (17.64 KB)
Non-trainable params: 0 (0.00 Byte)

The final complete Keras Model has the dictionary of `Input` objects as input and the classification result as outputs:

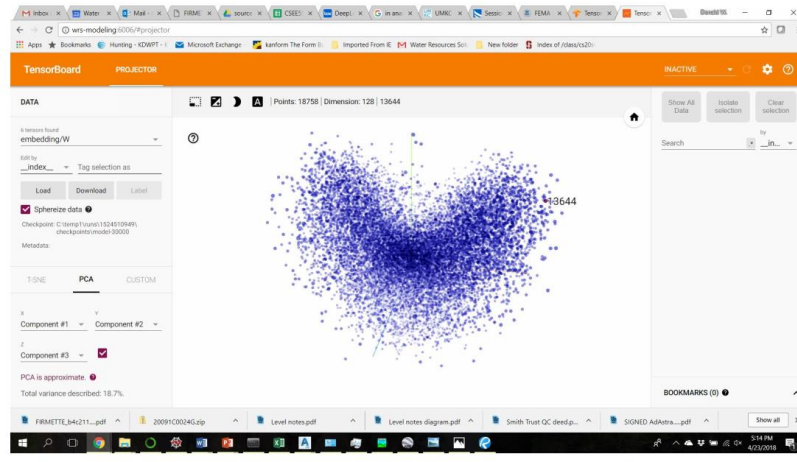
```
wage_model = tf.keras.Model(inputs, class_results, name="wage_model")
wage_model.summary()
```

```
Model: "wage_model"
=====
Layer (type)          Output Shape   Param #  Connected to
=====
age (InputLayer)      [(None, 1)]    0         []
education (InputLayer) [(None, 1)]    0         []
health (InputLayer)   [(None, 1)]    0         []
health_ins (InputLayer) [(None, 1)]    0         []
jobclass (InputLayer) [(None, 1)]    0         []
maritl (InputLayer)   [(None, 1)]    0         []
race (InputLayer)     [(None, 1)]    0         []
year (InputLayer)     [(None, 1)]    0         []
preproc (Functional)  (None, 35)     3         ['age[0][0]', 'education[0][0]', 'health[0][0]', 'health_ins[0][0]', 'jobclass[0][0]', 'maritl[0][0]', 'race[0][0]', 'year[0][0]']
classification (Sequential) (None, 4)  4516     ['preproc[0][0]']
=====
Total params: 4519 (17.66 KB)
Trainable params: 4516 (17.64 KB)
Non-trainable params: 3 (16.00 Byte)
```

With the model defined, it can now be compiled. That is, the loss function, the optimizer, and any additional metrics are specified. This example uses cross-entropy as a loss function, the Adam optimizer, and the accuracy and KL-divergence as additional metrics to calculate and report. The Python code block below explicitly shows some of the hyper-parameters for the Adam optimizer for illustration, but they are left at their defaults. Note how these parameters are used in Equations 13.9 to 13.11.

```
wage_model.compile(
    loss=tf.keras.losses.SparseCategoricalCrossentropy(
        from_logits=False),
    optimizer=tf.keras.optimizers.Adam(
        learning_rate=0.001,
        beta_1 = 0.9,
        beta_2 = 0.999,
        epsilon = 1e-07),
    metrics=[
        tf.keras.metrics.SparseCategoricalAccuracy(),
        tf.keras.metrics.KLDivergence()])
# Note: Specifying from_logits=True for the loss function can save
# the softmax activation or the softmax layer at the bottom of the
# sequential classification model.
```

As indicated in the model summary output above, the model has eight distinct inputs in its InputLayer. Hence, when fitting the model, the training or validation data must be provided as a dictionary with eight corresponding entries of Numpy arrays.



https://commons.wikimedia.org/wiki/File:Tensorboard_1.jpg

Figure 13.12: Tensorboard visualization

The following Python code block sets this up, converting Pandas dataframe columns to Numpy arrays:

```
import numpy as np
wage_feature_dict = \
    {name: np.array(value) for \
        name, value in wage_features.items()}
```

TensorBoard is a tool to visualize neural network models and their training and validation data/performance, as shown in Figure 13.12. To use the TensorBoard visualization tool, the training and validation performance is written to a set of files during the training process. The following code block specifies the folder where the files are to be created, with a separate subfolder for each model that is fitted, named by date and time. A callback function is then created that will be passed to the model `fit()` function:

```
import datetime
# Construct the folder name
log_dir = "./tensorboard_logs/" + \
    datetime.datetime.now().strftime("%Y%m%d-%H%M%S")

# Define the Tensorboard callback function
tensorboard_callback = \
    tf.keras.callbacks.TensorBoard(log_dir=log_dir, histogram_freq=0)
```

Next, the model is trained for 25 epochs, using 1/3 of the data for validation and the remainder for training:

```
wage_hist = wage_model.fit(
    x = wage_feature_dict,
    y = wage_labels,
    validation_split=0.333,
    batch_size=20,
    epochs = 25,
    callbacks=[tensorboard_callback])
```

Training history can be plotted using Python graphics packages, here Plotly Express:

```
import plotly.express as px
hist = pd.DataFrame({
    'training':
        wage_hist.history['sparse_categorical_accuracy'],
    'validation':
        wage_hist.history['val_sparse_categorical_accuracy']})
hist['epoch'] = np.arange(hist.shape[0])
hist = pd.melt(hist, id_vars='epoch',
               value_vars=['training', 'validation'])
fig = px.line(hist, x='epoch', y='value', color='variable')
fig.show()
```

However, TensorBoard provides a better and more interactive way to explore the model and its training performance. To use Tensorboard for models fitted with an appropriate callback function, first start Tensorboard from a terminal application or other bash shell and provide the log file directory that was specified in the TensorBoard callback definition:

This following command is NOT a Python command, use a bash shell in a Terminal!

```
tensorboard --logdir tensorboard_logs
```

TensorBoard is a web-based application, visit the URL <http://localhost:6006> in a web browser. Figures 13.13 to 13.15 show some excerpts of the TensorBoard visualizations. Figure 13.13 shows a summary of the training and validation performance, Figure 13.14 shows part of the computation graph that Tensorflow uses, showing the lookup layers defined in the model and the concatenate operation of the preprocessed inputs. Figure 13.15 shows an excerpt of the classification model, showing the dense and dropout layers.

Callback functions like the Tensorboard callback used above, are a Keras mechanism to perform certain actions as part of the training process. Keras calls these callback functions while training the model. Another useful callback function is the EarlyStopping callback. It allows the analyst to specify that Keras should stop

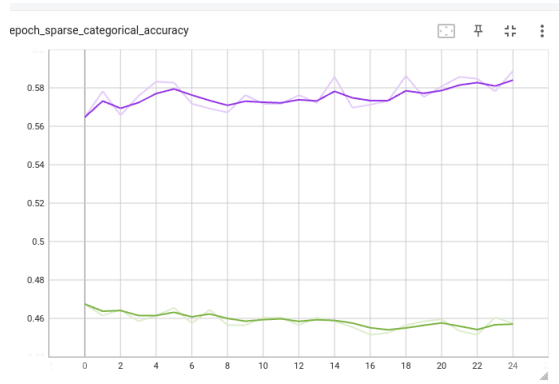


Figure 13.13: Tensorboard visualization of classification model (1)

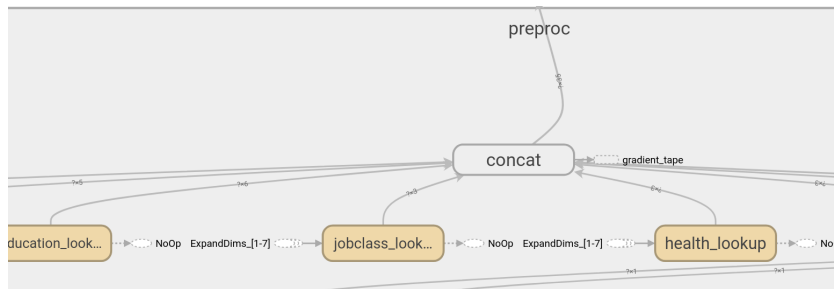


Figure 13.14: Tensorboard visualization of classification model (2)

model training when a certain metric does not change over a number of epochs. The following Python code block gives an example:

```
earlystop_callback = tf.keras.callbacks.EarlyStopping(
    monitor = 'val_loss',
    patience = 3,
    mode = 'min', # or 'max' or 'auto' depending on monitor metric
    restore_best_weights = True)
```

The metric to monitor can be the loss function (training or validation as per the prefix) or any of the additional metrics to be computed that are specified in the `metrics` argument to `compile()`. The `patience` parameter indicates the number of epochs to wait for an improvement before stopping the training; the `mode` parameter indicates whether to wait for decrease, increase, or to automatically determine this from the metric to monitor. Finally the `restore_best_weights` option is used to restore the parameter values from the best epoch or to retain the parameter values from the final training epoch. The early stopping callback can be provided to the `fit()` function in the same list as the Tensorboard callback.

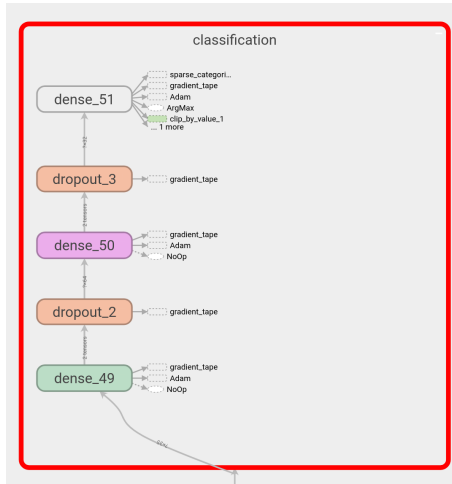


Figure 13.15: Tensorboard visualization of classification model (3)

Hands-On Exercise

- Examine the model summaries for the preprocessing, the classification, and the complete wage model. Explain the number of trainable and total parameters, and also explain the output shapes of each layer.
- Make the "wage" prediction a binary classification problem:
 1. Modify the `wage_labels` in the Pandas data frame and combine classes 0, 1 and classes 2, 3 (class numbers should be 0 or 1)
 2. Modify the classification network to have a single output node
 3. Use the `BinaryCrossentropy` loss
 4. Return the following metrics as part of the training history:
 - Precision
 - Recall
 - AUC
 5. Plot the metrics after training

13.7 Review Questions

Introduction

1. What is the original motivation behind the development of artificial neural networks (ANNs)?
2. Explain the role of biological neurons and their connections in the brain. How does this biological inspiration relate to ANNs?

Artificial Neural Networks

3. Describe the basic equation that defines a neural network unit or cell. Include the role of inputs, weights, bias, and the activation function.
4. Why is it important for the activation function to be non-linear in neural networks?
5. List at least three frequently-used activation functions and provide their mathematical definitions. Sketch them and their gradients.
6. What is the primary purpose of an activation function in a neural network?
7. Sketch and explain the architecture of a neural network with a single, fully-connected hidden layer.
8. Sketch a diagram of a non-trivial neural network and calculate the number of trainable parameters.
9. In the context of neural networks, what is the purpose of having an output layer with multiple output units?
10. How do neural networks handle multi-objective regression problems? Provide an example of a scenario where this might be useful.
11. Define multi-class classification in neural networks. How is the softmax activation function used in this context?
12. Write the mathematical expression for the softmax activation function and explain its components.
13. How does the architecture of a neural network influence its ability to learn from data? Consider the depth (number of layers) and width (number of units per layer) in your answer.

Parameter Estimation

14. What is the purpose of a loss function in training a neural network?
15. What loss functions are typically used for classification in neural networks?
16. Define the parameter vector ϕ of a neural network. What are its components?
17. How does the choice of loss function affect the training and performance of a neural network?
18. Compare and contrast the Mean Squared Error (MSE) and Mean Absolute Error (MAE) functions. When might you choose one over the other?

Gradient Descent

19. What is the gradient of a function and how is it used in gradient descent?

20. Explain the iterative process of gradient descent. What are the steps involved?
21. Describe the potential problems encountered during gradient descent.
22. Why is it important to choose an appropriate learning rate in gradient descent?

Stochastic Gradient Descent

23. How does stochastic gradient descent (SGD) differ from non-stochastic gradient descent? Why is it called "stochastic"?
24. What is a minibatch in the context of SGD, and why are minibatches used?
25. Define an epoch in the context of training neural networks.
26. Explain the concept of early stopping and its purpose in training neural networks.
27. How does the size of a minibatch affect the performance and stability of SGD?
28. What are the advantages and disadvantages of using large versus small mini-batches in SGD?

Parameter Updates

29. What is an adaptive learning rate and why is it used?
30. Describe the AdaGrad method and its purpose.
31. How does RMSProp modify AdaGrad to improve its performance?
32. Explain the AdaDelta method and how it differs from RMSProp.
33. What are momentum methods in SGD and what problem do they address?
34. Describe the Nesterov momentum method and how it improves upon standard momentum.

Gradient Problems

35. What are vanishing gradients and why do they occur in neural networks?
36. List and explain three methods to address the vanishing gradient problem.
37. Describe the ResNet architecture and how it helps to address the vanishing gradients problem.
38. Explain the concept of gradient clipping and its purpose.
39. How does batch normalization help address the vanishing gradient problem?
40. Discuss the importance of proper weight initialization in preventing gradient problems.

Regularization with Dropout

41. What is dropout and how does it help to prevent overfitting in neural networks?
42. What are typical dropout rates used in practice?
43. Explain how dropout mimics the training of an ensemble of neural networks.
44. Discuss the impact of dropout on the training and evaluation phases of neural network training.
45. How does dropout act as a form of regularization in neural networks?

Chapter 14

Convolutional Neural Networks

Sources and Further Reading

The material in this chapter is based on the following sources.

Kevin P. Murphy: *Probabilistic Machine Learning – An Introduction*. MIT Press 2022.

<https://probml.github.io/pml-book/book1.html>

Chapter 13, 14, 15

The book by Murphy is freely available and provides three chapters on neural networks, one for structured data, one for images, and one for sequences. It provides significant depth on convolutional and recurrent network architectures, fitting the models, and problems that the data analyst may encounter.

Tensorflow Guides:

<https://www.tensorflow.org/guide>

https://www.tensorflow.org/tutorials/load_data/text

This course uses the Tensorflow programming framework for neural network applications. The Tensorflow website has a multitude of introductory and advanced guides and tutorial that cover all aspects of machine learning with neural networks.

14.1 Introduction

Convolutional Neural Networks (CNNs) are a class of deep neural networks, most commonly applied to analyzing images. They are also known as *ConvNets* and are particularly powerful for tasks like image recognition and classification. CNNs have been successful in identifying faces, objects, and traffic signs as well as powering vision in robots and self driving cars. CNNs as they are known today were popularized in the 1990s by Yann LeCun and his colleagues, who used them for digit recognition in postal codes.

The primary motivation for convolutional networks stems from the limitations of fully connected networks in image processing. Fully connected networks that treat input pixels independently lack the spatial hierarchies of features in an image, which leads to inefficiencies and a loss of spatial relationships among pixels. CNNs address these issues by leveraging spatial hierarchies through localized receptive fields called *convolution filters*, shared weights, and pooling, which results in robustness to image transformations and significant reduction in the number of parameters needed.

At its core, a CNN automatically learns and identifies various features in images at multiple levels of abstraction. For example, the initial layers might detect edges and textures, intermediate layers learn to identify larger motifs, and deeper layers interpret these motifs as parts of familiar objects. CNNs have been successfully applied in a wide range of visual data applications, including:

- *Image and Video Recognition*: CNNs can classify images and videos into categories, often surpassing human performance in tasks such as facial recognition and object detection.
- *Medical Image Analysis*: In healthcare, CNNs are used for diagnosing diseases by analyzing medical scans to detect anomalies like tumors in MRI scans or X-rays.
- *Autonomous Vehicles*: They help in identifying obstacles, understanding traffic signs, and enabling vehicles to make informed decisions.
- *Augmented Reality*: CNNs can augment real-world environments by enhancing image and video feeds in real-time, providing a richer user experience.

CNNs represent a significant advancement in the ability to automatically interpret large sets of visual data, leading to improvements across various applications that require automatic visual recognition. Their ability to understand the complexity of images and videos with increasing accuracy has made them the backbone of modern artificial intelligence in the visual domain.

14.2 Convolutional Layers

A *convolutional layer* is a fundamental building block of a Convolutional Neural Network (ConvNet). It performs the primary operations that allow a ConvNet to capitalize

on the spatial structure of input data, such as images, by extracting features that become increasingly complex and high-level as data progresses through deeper layers of the network.

A convolutional layer applies a set of learnable *convolution filters* (also known as *convolution kernels*) to the input. Each filter is small spatially (along width and height), but extends through the full depth of the input volume.

The convolution operation involves sliding each filter across the width and height of the input image and computing dot products between the filter and the local regions of the input image. As the filter slides over the input image, a 2-dimensional activation map (or feature map) is created as output that gives the responses of that filter at every spatial position. Intuitively, the network learns filters that activate when they see some specific type of feature at some spatial position in the input.

Figure 14.1 shows an example in one dimension. The kernel vector $[1, 2]$ is first multiplied element-wise with the first two elements of the input, yielding $[0, 1][1, 2] = [0, 2]$. The result is then summed, yielding 2. This is the first element of the output. The kernel is then moved one position to the right. Multiplication with that input portion yields $[1, 2][1, 2] = [1, 4]$ and summing yields 5, the second element of the output or activation/feature map. This is done until the kernel is multiplied with the two right-most elements of the input in Figure 14.1. Note that the kernel of length 2 can be applied 6 times to the input of length 7 in Figure 14.1, yielding an output of length 6.

Figure 14.2 shows an example in two dimensions. Here, the first multiplication and summing yields $0\hat{0} + 1\hat{1} + 3\hat{2} + 4\hat{3} = 19$, which is the top left element of the output in Figure 14.2. The kernel can then be moved once to the right, and once to the bottom. This yields an output shape of 2×2 .

Figure 14.3 shows a multi-channel 2D convolution. The kernel consists of two channels that are applied to two channels of the input. There are as many channels in the kernel as there are channels in the input. The two channels are then summed to produce a single output channel. As a realistic example, consider an RGB image with three "depth" channels, one each for the red, green, and blue color components. A kernel might then have a size of $3 \times 3 \times 3$ (3 width, 3 height, and 3 for the three color channels).

A single convolutional network layer may have *multiple filters*. Each filter will yield one output channel or activation/feature map. These output channels are in addition to the depth of each filter. For example, the $2 \times 2 \times 2$ filter applied to the $3 \times 3 \times 2$ input in Figure 14.3 yielded a single output channel of shape 2×2 . Applying 10 independent filters that are each of shape $2 \times 2 \times 2$ to the input yields an output of shape $2 \times 2 \times 10$.

Input	Kernel	Output	
$\begin{bmatrix} 0 & 1 & 2 & 3 & 4 & 5 & 6 \end{bmatrix}$	$\begin{bmatrix} 1 & 2 \end{bmatrix}$	$=$	$\begin{bmatrix} 2 & 5 & 8 & 11 & 14 & 17 \end{bmatrix}$

Source: Murphy Fig. 14.4

Figure 14.1: 1-dimensional convolution filter

Input	Kernel	Output
$\begin{bmatrix} 0 & 1 & 2 \\ 3 & 4 & 5 \\ 6 & 7 & 8 \end{bmatrix}$	$\begin{bmatrix} 0 & 1 \\ 2 & 3 \end{bmatrix}$	$= \begin{bmatrix} 19 & 25 \\ 37 & 43 \end{bmatrix}$

Source: Murphy Fig. 14.5

Figure 14.2: 2-dimensional convolution filter

Consider another simple example in image processing, with an input image of size $32 \times 32 \times 3$ (width x height x channel (red, green, blue)). A convolutional layer might have 10 filters of size $5 \times 5 \times 3$. The output of applying these filters will result in an output of size $28 \times 28 \times 10$ (when using a stride of 1 and no padding, see below).

While the examples show fixed numbers in the kernels, the kernel elements are actually trainable parameters.

Verifying the shape/dimensions of inputs, kernels, and outputs, and understanding how they are related to each other is important to understand and to verify the correctness of the implemented model!

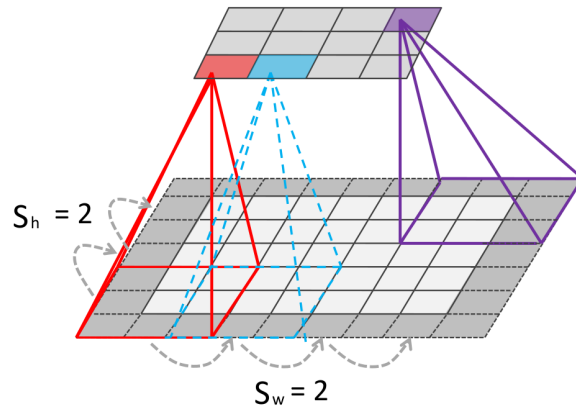
There are two key considerations when defining a convolutional network layer, *stride* and *padding*. When moving the kernel over the input, the *stride* controls the size of the steps in each direction. A stride of 1 means the kernel moves one column or row at a time. A larger stride results in downsizing of the feature map, in removing redundancy in the output feature map (it avoids an input element being processed multiple times by a filter) and in increasing the efficiency of the data representation in the neural network.

Second, the input is often *padded* with zeros around the edges ("zero-padding"). This allows control over the shape of the output feature map and it also increases the weight

Input	Kernel	Input	Kernel	Output
$\begin{bmatrix} 0 & 1 & 2 & 3 \\ 3 & 4 & 5 & 6 \\ 6 & 7 & 8 & 9 \end{bmatrix}$	$\begin{bmatrix} 0 & 1 & 2 \\ 3 & 4 & 5 \\ 6 & 7 & 8 \end{bmatrix}$	$\begin{bmatrix} 1 & 2 & 3 \\ 4 & 5 & 6 \\ 7 & 8 & 9 \end{bmatrix}$	$\begin{bmatrix} 1 & 2 \\ 3 & 4 \end{bmatrix}$	$= \begin{bmatrix} 56 & 72 \\ 104 & 120 \end{bmatrix}$
			$\begin{bmatrix} 0 & 1 \\ 2 & 3 \end{bmatrix}$	$+ = \begin{bmatrix} 56 & 72 \\ 104 & 120 \end{bmatrix}$

Source: Murphy Fig. 14.9

Figure 14.3: Multi-channel 2D convolution



Source: Murphy Fig. 14.8(b)

Figure 14.4: Striding and padding in a convolutional network

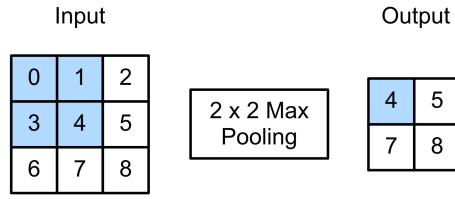
or emphasis given to edge values, which are otherwise captured only once by a filter, in contrast to interior values, which are captured multiple times as the kernel moves over the input. Inputs may be padded by one "frame" of zeros or by multiple zero-frames, limited only by the size of the convolution kernel.

Figure 14.4 shows an example of a zero padded input (bottom, dark gray squares surrounding the light gray squares indicate padding) and a striding of 2 in each direction. The output feature map is shown on top.

While the convolution layer may look complicated and complex, it essentially produces a weighted sum of inputs where the convolution kernel entries are the weights, very much like the general neural network unit definition in the previous chapter. A bias term is added to the weighted sum produced by the convolution, before the sum is transformed using a non-linear activation function. In many ConvNets, the activation function is the ReLU (Rectified Linear Unit). In other words, the CNN functions very much like the general neural networks introduced in the previous chapter, except that the connections between inputs and weights are more complex and focus on spatial relationships. As with fully connected networks, it is important to understand the number of parameters (weights and biases) in order to verify the correctness of the implemented model.

14.3 Pooling Layers

Pooling layers within ConvNets are used to reduce the spatial dimensions (width and height) of the convolution output for the next convolutional layers or for subsequent dense, fully-connected layers. This reduction in dimensions serves several purposes. It decreases the computational load and the memory usage, helps prevent overfitting by providing an abstracted form of the representation, and makes the network invariant



Source: Murphy Fig. 14.12

Figure 14.5: Pooling in a ConvNet

to small transformations, distortions, and translations in the input. Pooling is applied separately to each output channel of a convolution.

The most common types of pooling include *max pooling* and *average pooling*. Max pooling partitions the convolution output into a set of overlapping or non-overlapping regions, and outputs the maximum value for each such region. For instance, a 2×2 max-pooling layer applied with a stride of 2 reduces the size of the input by a factor of four (halving the height and width separately). Instead of taking the maximum value from each distinct region, average pooling computes the average of the values in each patch on a feature map.

Figure 14.5 shows an example of 2×2 max pooling on a 3×3 input with stride 1. In this example, the pooling stride is less than the pooling filter size, so that the pooled regions are overlapping. Each convolution output may be captured multiple times in different pooling outputs. When the stride is chosen to the same size as the pooling filter, the regions are not overlapping. Overlapping regions retain more information but are computationally more intensive and require additional memory, as the size of the pooling output and therefore the size of all subsequent layers is larger.

In image processing in particular, pooling helps the network become invariant to small translations, rotations, and scalings of an image. Max pooling in particular helps in detecting features like edges, textures, and shapes, even if they appear in different areas of the image in different instances. Although pooling reduces the resolution of the feature map, important features (like the presence of certain edges or textures) are retained due to the nature of max or average operations within localized regions of the input.

Hands-On Exercise

1. Assume the following 5×5 input matrix:

$$\begin{bmatrix} 1 & 2 & 3 & 2 & 1 \\ 2 & 3 & 2 & 1 & 3 \\ 3 & 2 & 1 & 3 & 2 \\ 2 & 1 & 3 & 2 & 1 \\ 1 & 3 & 2 & 1 & 3 \end{bmatrix}$$

and the following 3×3 convolution kernel:

$$\begin{bmatrix} 1 & 2 & 1 \\ 2 & 4 & 2 \\ 1 & 2 & 1 \end{bmatrix}$$

Zero-pad the matrix with *two* zeros on all sides and using stride 2, calculate the convolution result. What are the dimensions of the convolution output?

2. Apply a 2×2 max pooling layer to the result of the previous exercise.

14.4 Understanding ConvNets

The convolutional and pooling layers of a CNN are only part of the complete neural network. The purpose of the these layers is essentially *feature extraction* or *feature learning*, that is, to identify or learn features in the input that might be useful for subsequent tasks like classification. For images processing tasks, the convolutional and pooling layers may detect features such as edges, textures, geometric shapes, etc. These features are useful in classifying the image content, e.g. does the image show a bee or a tiger.

Figure 14.6 shows an example of a complete network. Multiple convolutional and pooling layers are stacked in a sequential model. The output of the final pooling layer is then "flattened", that is, the information in the two or three dimensional output (width x height x channels) is arranged linearly in a one-dimensional vector. This vector forms the input to a classification network, which typically consists of a sequence of dense, fully-connected layers and a final softmax output layer, as illustrated in the previous chapter. Thus, one can think of the convolutional part of the network as the feature extraction or feature learning part whose output are the features for the classification part of the network.

Figure 14.7 shows another useful diagram. In addition to the general architecture it shows the dimensionality of the output at each stage, which is very helpful in understanding how convolution and pooling filters operate. For example, the input image is

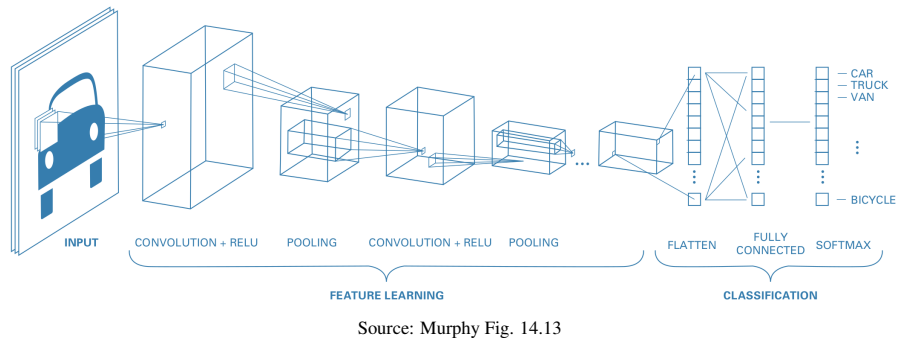
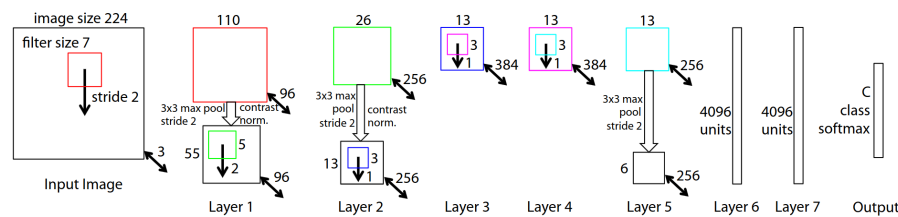


Figure 14.6: Convolutional network for image classification



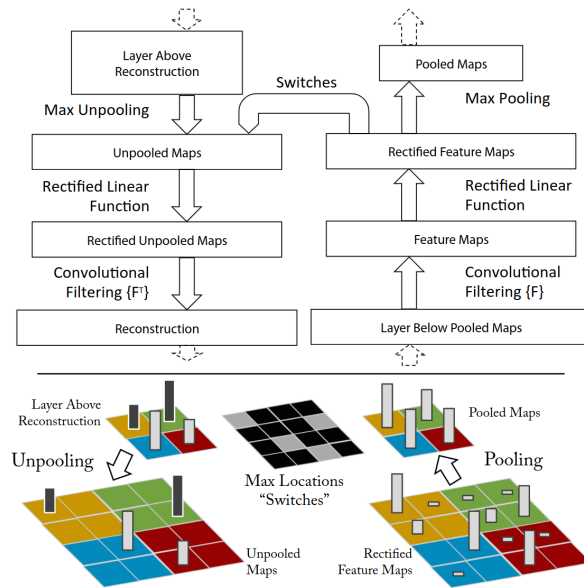
Source: Matthew D. Zeiler and Rob Fergus (2013) Visualizing and Understanding Convolutional Networks.
<https://doi.org/10.48550/arXiv.1311.2901>

Figure 14.7: Convolutional network for image classification, with dimensionality

of shape $224 \times 224 \times 3$. There are 96 convolutional filters operating on this image, each with a kernel shape of $7 \times 7 \times 3$ and a stride of 2. This yields an output with shape $110 \times 110 \times 96$. The subsequent 3×3 pooling with stride 2 reduces this to $55 \times 55 \times 96$ for the final layer 1 output.

Verifying the shape/dimensions of inputs, kernels, and outputs, and understanding how they are related to each other is important to understand and to verify the correctness of the implemented model!

This term "feature map" may be used in various ways. Some authors use it to describe the convolution output before the application of activation function and pooling, some authors use it to describe the convolution output after application of the activation function but prior to pooling, some authors use it to describe the output after pooling and some authors use it to describe any of the three. Hence, when the term "feature map" is used, it is important to identify the precise meaning in a particular context.



Source: Matthew D. Zeiler and Rob Fergus (2013) Visualizing and Understanding Convolutional Networks.
<https://doi.org/10.48550/arXiv.1311.2901>

Figure 14.8: DeconvNet architecture

To understand how convolutional networks perform feature extraction or feature learning, it is instructive to visualize the feature maps. There are multiple ways of doing this. One of the earliest techniques developed for this is the "*DeconvNet*¹". The general architecture of a DeconvNet is shown in Figure 14.8. Using an unpooling operation, a DeconvNet successively reconstructs the image input for a feature map.

Figures 14.9 to 14.12 show reconstructed visualizations for a selection of feature maps in a 5-layer convolutional network. The feature maps on the left of each image give an indication of the type of feature they represent, and the corresponding visualization on the right of each figure gives examples of input images that correspond to the detected features. The images on the right of each figure are not training images, they are reconstructed by the DeconvNet.

Figure 14.9 indicates that the first layer appears to have learned to recognize basic features such as edges or image contrasts as well as colours of an image.

The second layer in Figure 14.10 appears to have learned to recognize geometric shapes such as lines, circles, squares, but also colors.

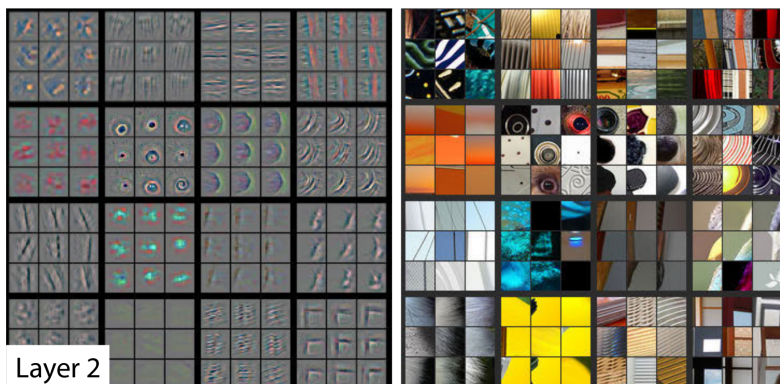
Layer 3, shown in Figure 14.11, has learned to recognize more complex features that

¹Matthew D. Zeiler and Rob Fergus (2013) Visualizing and Understanding Convolutional Networks.
<https://doi.org/10.48550/arXiv.1311.2901>



Source: Matthew D. Zeiler and Rob Fergus (2013) Visualizing and Understanding Convolutional Networks.
<https://doi.org/10.48550/arXiv.1311.2901>

Figure 14.9: Layer 1 feature map visualization by a DeconvNet

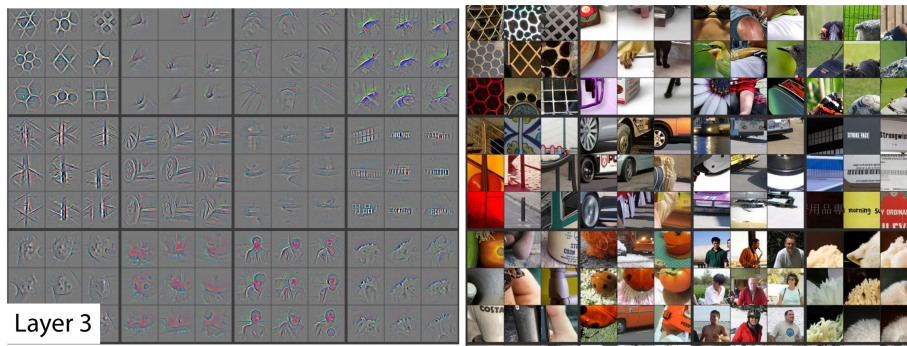


Source: Matthew D. Zeiler and Rob Fergus (2013) Visualizing and Understanding Convolutional Networks.
<https://doi.org/10.48550/arXiv.1311.2901>

Figure 14.10: Layer 2 feature map visualization by a DeconvNet

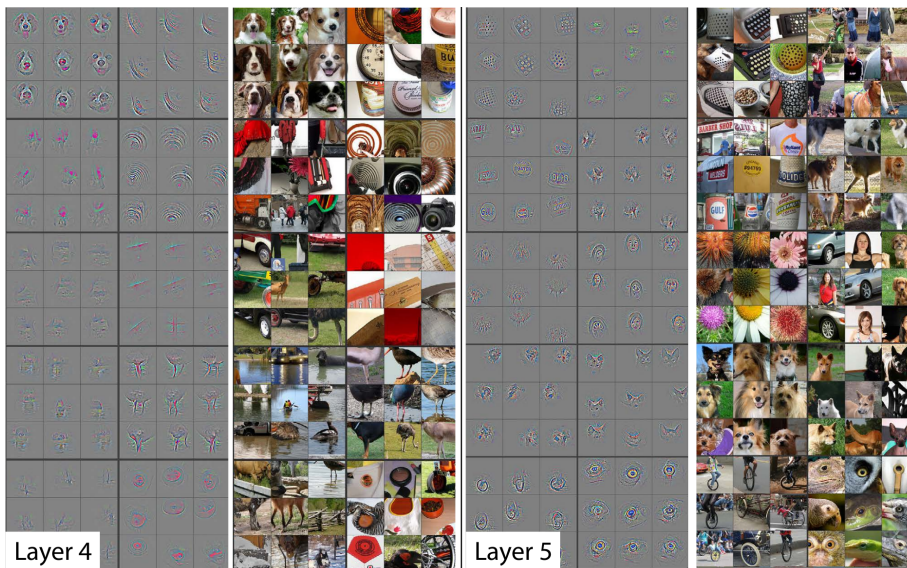
resemble simple objects, such as wheels, fruits, people, but also complex geometric patterns.

Finally, layers 4 and 5 have learned to extract specific objects or object parts, such as dogs, birds, but also geometric shapes. The final layer shows a clear ability to extract the presence of specific object classes in an image.



Source: Matthew D. Zeiler and Rob Fergus (2013) Visualizing and Understanding Convolutional Networks.
<https://doi.org/10.48550/arXiv.1311.2901>

Figure 14.11: Layer 3 feature map visualization by a DeconvNet



Source: Matthew D. Zeiler and Rob Fergus (2013) Visualizing and Understanding Convolutional Networks.
<https://doi.org/10.48550/arXiv.1311.2901>

Figure 14.12: Layers 4 and 5 feature map visualization by a DeconvNet

14.5 Image Classification Example using Tensorflow

This section illustrates how to implement convolutional neural networks in Tensorflow/Keras². This example uses the CIFAR10 data set, a standard data set used to

²Python code for this example is taken from TensorFlow.org that is made available under a [Apache 2.0 license](#).

benchmark image classification methods. The data set consists of 60,000 images with resolution of 32×32 pixels and 3 RGB channels. There are ten classes. The latest error rates on this data set are less than 0.5%, showing that it is no longer useful as a realistic or competitive benchmark — it is essentially a solved problem. However, due to the small image size and data set size, it is computationally easy to analyze and useful as an illustrative example.

Complete implementations for this and the other examples in this chapter are available in the following GitHub repo: <https://github.com/jevermann/busi4720-ml>

The project can be cloned from this URL: <https://github.com/jevermann/busi4720-ml.git>

The data set is part of the Keras Python package, making import easy. The function `load_data()` provides separated training and testing images and labels:

```
import tensorflow as tf
from tensorflow.keras import datasets, layers, models
import datetime

(train_images, train_labels), (test_images, test_labels) \
    = datasets.cifar10.load_data()
```

The RGB values in the original images are coded on a scale of 0 to 255. The first step is to scale these values to the range of 0 to 1 in order to avoid gradient problems during model training:

```
# Normalize pixel values to be between 0 and 1
train_images, test_images = train_images/255.0, test_images/255.0
```

Keras makes it easy to create a convolutional network similar to the one shown in Figure 14.7. This example uses three convolutional layers, each with a ReLU activation and followed by max-pooling.

The first argument to the `Conv2D()` layer creation function specifies the number of output channels, that is, the number of independent convolution kernels. In the example, the first layer has 8 output channels, the second and third layers have 16 output channels. The second argument specifies the shape of the convolution kernel, which is 3×3 for all layers in this example. The input shape for the first convolutional layer must be provided explicitly. Note that no batch size is specified, only the image input dimensions are needed; the batch size can be specified later. The input shapes for subsequent layers are automatically determined from the output shapes of the previous layers.

```
# Create a simple convolutional model:
model = models.Sequential()
model.add(layers.Conv2D(8, (3, 3), activation='relu',
                      input_shape=(32, 32, 3)))
model.add(layers.MaxPooling2D((2, 2)))
model.add(layers.Conv2D(16, (3, 3), activation='relu'))
model.add(layers.MaxPooling2D((2, 2)))
model.add(layers.Conv2D(16, (3, 3), activation='relu'))
```

After the convolutional layers for feature extraction, the remainder of network uses fully-connected, dense layers for classification. Because there are 10 target classes, the final, output layer has 10 units.

```
# Add dense (fully-connected) layers for classification.
# There are 10 target classes
model.add(layers.Flatten())
model.add(layers.Dense(32, activation='relu'))
model.add(layers.Dense(10))
# Show model summary
model.summary()
```

The model summary is shown below. It is easy to verify the number of parameters. For example, the first Conv2D layer uses 3×3 kernels on 3 channels, for a total of 27 parameters. There are 8 such kernels, each creating one of the 8 output channels, and each adding a bias term before applying the activation function, for a total of $8 \times 27 + 8 = 224$ parameters. The output shape is 30×30 because a kernel of shape 3×3 can be moved 30 times in each direction with a stride of 1.

The max pooling layer uses a kernel size of 2×2 and the default for Keras max-pooling layers is to use non-overlapping regions, that is, a stride of the same size as the kernel size. Hence, the output shape of the first max-pooling layer is 15×15 for each of the 8 channels.

The second Conv2D layer operates on this input size and produces 16 output channels with a kernel size of 3×3 . It needs to operate on the 8 channels of the input yielding $3 \times 3 \times 8 = 72$ parameters for each kernel. There are 16 of such kernels that each create one output channel. Each such kernel again adds one bias term to the weighted sum of the convolution for a total of $72 \times 16 + 16 = 1168$ parameters. With a stride of 1, the kernel can be moved across the input 13 times, resulting in an output shape of $13 \times 13 \times 16$. The secon max-pooling layer also uses a 2×2 non-overlapping kernel, yielding an output size of $6 \times 6 \times 16$.

The third Conv2D layer uses 16 output channels of a 3×3 kernel. Each kernel operates on 16 input channels. Thus, the total number of parameters in this layer is $3 \times 3 \times 16 \times 16 + 16 = 2320$. A kernel this size can be moved 4 times in each direction over an input of size 6×6 , yielding a final output shape of $4 \times 4 \times 16$ for a total of 256 values. The flatten layer simply transforms them to a one-dimensional layer of 256 values.

The following dense layer with 32 units then has $32 \times 256 + 32 = 8224$ trainable parameters (refer to the previous chapter for details), and the output layer for 10 classes has $10 \times 32 + 10$ parameters. Note that the model contains no softmax activation or softmax layer, so the final output are logits, not class membership probabilities.

Layer (type)	Output Shape	Param #
<hr/>		
conv2d (Conv2D)	(None, 30, 30, 8)	224
max_pooling2d (MaxPooling2D)	(None, 15, 15, 8)	0
conv2d_1 (Conv2D)	(None, 13, 13, 16)	1168
max_pooling2d_1 (MaxPooling2D)	(None, 6, 6, 16)	0
conv2d_2 (Conv2D)	(None, 4, 4, 16)	2320
flatten (Flatten)	(None, 256)	0
dense_3 (Dense)	(None, 32)	8224
dense_4 (Dense)	(None, 10)	330
<hr/>		
Total params: 12266 (47.91 KB)		
Trainable params: 12266 (47.91 KB)		
Non-trainable params: 0 (0.00 Byte)		

Verifying the shape/dimensions of inputs, kernels, and outputs, and understanding how they are related to each other is important to understand and to verify the correctness of the implemented model!

Next, the callback function for writing TensorBoard log files is specified, similar to examples in the previous chapter:

```
# Construct log file folder name
log_dir = "./tensorboard_logs/" + \
    datetime.datetime.now().strftime("%Y%m%d-%H%M%S")
# Create TensorBoard callback function
tensorboard_callback = \
    tf.keras.callbacks.TensorBoard(log_dir=log_dir, histogram_freq=0)
```

Finally, the model is compiled and trained for 100 episodes. Note that `from_logits` must be set for the loss function because the model's output layer provides logits, not probabilities.

```
# Compile and train the model:
model.compile(optimizer='adam',
    loss=tf.keras.losses \
        .SparseCategoricalCrossentropy(from_logits=True),
    metrics=['accuracy'])

history = model.fit(train_images, train_labels,
    epochs=10,
    validation_data=(test_images, test_labels),
    callbacks=[tensorboard_callback])
```

The training accuracy after 100 episodes is $\approx 75\%$ and the validation accuracy is $\approx 66\%$ showing relatively poor performance and some evidence of overfitting. However, noting that random performance should yield an accuracy of 0.1 for 10 classes, it also shows that a relatively simple network with only a few thousand parameters is already quite capable at image classification.

Hands-On Exercise

Adapt the network architecture to identify the impact on training and validation performance of the following:

1. Convolution kernel size (originally 3×3)
2. Number of convolution filters (output channels) (originally 8, 16)
3. Convolution stride (originally 1)
4. Number of 2D-Conv layers (originally 3)
5. Type of pooling (originally max-pooling)

Comment on your findings and identify the best model.

Tip: You can find information on how to change the layers in the Keras documentation for [convolution layers](#) and for [pooling layers](#).

In particular, explore the trade-offs between using a large number of smaller filters versus a smaller number of larger filters in the context of convolution followed by pooling.

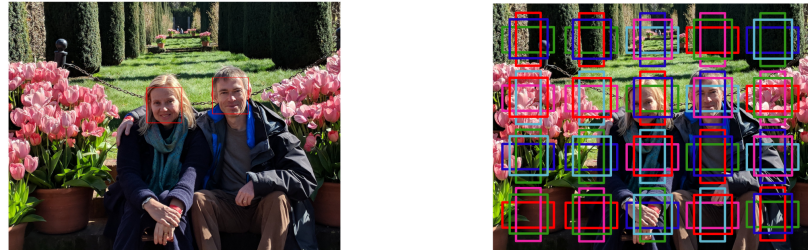
Explore how does the choice of pooling strategy (max vs average) affects the features extracted by a convolutional neural network and the predictive performance of the model.

14.6 Other Computer Vision Tasks for CNNs

Two additional tasks that CNNs are useful for are object detection and semantic segmentation. These two tasks are the basis of a wide array of applications including autonomous driving, medical image analysis, and surveillance.

Object Detection is the task of identifying and localizing objects within an image or a video. The task is not just to classify whether certain objects are present (as in image classification) but also to find their boundaries or "bounding boxes" within the image. Object detection is crucial in scenarios where the context provided by the location of an object within a scene affects the decision-making process, such as in security for detecting suspicious activities or in retail for counting products.

Figure 14.13 illustrates the problem for the input image in the left panel. The right panel overlays this input image with a set of potential bounding boxes or "anchor boxes". For each anchor box, the CNN must learn the object presence probability,



Source: Murphy Fig. 14.27

Figure 14.13: Object detection example input and bounding boxes

the object category, and two offset vectors that to be added to the center of the box that shifts the position of the box and scales the size of the box. Formally, the problem is defined as function f_θ of some parameters θ such that:

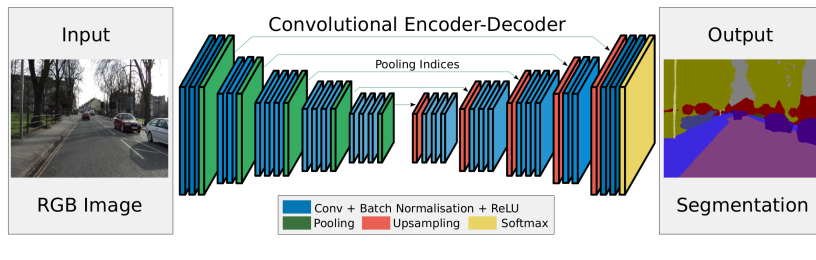
$$f_\theta : R^{H \times W \times K} \rightarrow [0, 1]^{A \times A} \times \{1, \dots, C\}^{A \times A} \times (R^4)^{A \times A}$$

Early approaches like R-CNN (Regions with CNN features) generated potential bounding boxes in an image and then ran a classifier on these regions to detect objects. More advanced models like YOLO (You Only Look Once) and SSD (Single Shot Detector) view this as a single regression problem, straight from image pixels to bounding box coordinates and class probabilities.

Semantic Segmentation is the task of classifying each pixel of an image into a pre-defined category. This is more granular than object detection, which only discerns objects at the bounding box level. Semantic segmentation is used extensively in medical imaging (e.g., to delineate different types of tissue), road segmentation for autonomous vehicles, and landscape classification in geographic information systems.

Figure 14.14 shows an example input image on the left. The task is to identify the road surface, sidewalks, background objects, and the sky. An example output is shown on the right of that figure with different colors indicating the different classes of image pixels. One way to tackle the problem is with a convolutional encoder-decoder network as shown in the center of the figure.

For some applications, object detection and semantic segmentation must be combined. Consider as an example the input image for an autonomous vehicle task shown in Figure 14.15. Object detection delivers bounding boxes and image classes while subsequent semantic segmentation within each bounding box can provide the specific shapes of the objects.



Source: Murphy Fig. 14.29

Figure 14.14: Semantic segmentation example input, network architecture, and output



https://github.com/matterport/Mask_RCNN

Figure 14.15: Object detection and semantic segmentation for an autonomous driving task

The Keras web site has a large number of Keras tutorials on all kinds of computer vision problems, including object detection and semantic segmentation. See this site for more information: <https://keras.io/examples/vision/>.

14.7 Text Classification Example using Tensorflow

Convolutional neural networks, while originally developed and most widely used for image and video processing, have also been adapted to text classification tasks. This adaptation builds on the hierarchical structure of text, much like the hierarchical features in images, to effectively classify text into predefined categories. Common applications include sentiment analysis, topic labeling, spam detection, and others. CNNs have emerged as an alternative to RNNs (recurrent neural networks).

The two examples in this section use the StackOverflow 16k data set³. This data set

³Python code for this example is taken from TensorFlow.org that is made available under a [Apache 2.0 license](#).

consists of 16,000 user-generated questions about programming problems from the StackOverflow web site. The task is to classify the questions by programming language. Four classes are defined: CSharp, Java, JavaScript, and Python. The data set can be downloaded directly from the Tensorflow website.

First, import the required packages:

```
import collections
import pathlib
import datetime
import tensorflow as tf
from tensorflow.keras import layers, losses, utils
from tensorflow.keras.layers import TextVectorization
```

Next, download the StackOverflow data set. The data set will be placed in the local Keras cache directory and will be separated by training and test data, as well by the four classes, as shown in Figure 14.16.

```
# Get the data
url='http://download.tensorflow.org/data/stack_overflow_16k.tar.gz'
dataset_dir = utils.get_file(origin=url, untar=True,
    cache_subdir='stack_overflow')

# Remember where we put it
dataset_dir = pathlib.Path(dataset_dir).parent
train_dir = dataset_dir/'train'
test_dir = dataset_dir/'test'

# Print a sample
sample_file = train_dir/'python/1755.txt'
with open(sample_file) as f:
    print(f.read())
```

The sample output from the code above is shown here (training data for python, observation 1755). Note that the actual input has been "blinded", that is, direct mentions of the programming language have been removed.

```
why does this blank program print true x=true.def stupid()..
x=false.stupid().print x
```

Next, the training data portion is split into training and validation sets, yielding a total of three portions of data: training, validation, and testing:

✓ .keras		3 items	15:44	☆
> datasets		4 items	15:43	☆
✓ stack_overflow		4 items	15:44	☆
> test		4 items	26 May 2020	☆
> csharp		2,000 items	10 Jun 2020	☆
> java		2,000 items	10 Jun 2020	☆
> javascript		2,000 items	10 Jun 2020	☆
> python		2,000 items	10 Jun 2020	☆
> train		4 items	26 May 2020	☆
> csharp		2,000 items	10 Jun 2020	☆
> java		2,000 items	10 Jun 2020	☆
> javascript		2,000 items	10 Jun 2020	☆
> python		2,000 items	10 Jun 2020	☆
README.md		529 bytes	8 Jul 2020	☆
stack_overflow_16k.tar.gz		6.1 MB	15:44	☆

Figure 14.16: Screenshot of StackOverflow 16k data set downloaded to local Keras cache directory

```
raw_train_ds=utils.text_dataset_from_directory(
    train_dir, batch_size=32, validation_split=0.2,
    subset='training', seed=42)

raw_val_ds=utils.text_dataset_from_directory(
    train_dir, batch_size=32, validation_split=0.2,
    subset='validation', seed=42)

raw_test_ds=utils.text_dataset_from_directory(test_dir, batch_size=32)
```

The first step in applying neural networks to text data is to pre-process and standardize the text representation itself. This includes at least the following activities:

- *Standardization* converts text to lower-case, repairs spelling errors, removes stop words (such as "if", "the" etc.), removes punctuation (commas, periods), HTML code, and other special characters, and stems words (for example, turning "programming" into "program")
- *Tokenization* splits character strings into separate tokens, for example splitting sentences into words on whitespace (spaces, tab characters, etc.), or splitting words into subwords (for example, "subword" is split to "sub" and "word").
- *Vectorization* converts tokens into numerical values for input to the neural network. It uses methods such as one-hot encoding, word embeddings, or relative word frequencies.

14.7.1 Text classification using Bag-of-Word encoding without a CNN

This example uses the "Bag-of-Words" word frequency model that encodes the input as the number of times that a word occurs. Consider the following example where the input sentence is represented as a dictionary of word counts.

```
John likes to watch movies. Mary likes movies too.
{"John":1,"likes":2,"to":1,"watch":1, "movies":2,"Mary":1,"too":1}
```

This is a simple encoding that may suffice for the classification task, but it does neglect word order which could be important for other tasks. The second example below will use a different encoding that respects word order and is therefore suitable for the use with CNNs. This first, simple example serves as a baseline to compare the later example to.

Keras provides an easy way to preprocess text using the `TextVectorization` layer. The following Python code block creates such a layer for a maximum of 10,000 different tokens (approximately the first 10,000 unique words). It standardizes the input by converting all words to lower-case and removing punctuation. Tokenization then splits the input on whitespace into separate words. The words are not stemmed or processed further. Finally, vectorization produces "multi-hot" output, that is, the frequency of words for each input as seen in the above example. The `TextVectorization` layer is then adapted to the training set. Essentially, adaptation creates the list of 10,000 unique words to count when the actual input is processed.

```
# Use the Keras TextVectorization pre-processing layer:
multi_hot_vectorize_layer = TextVectorization(
    max_tokens=10000,
    standardize='lower_and_strip_punctuation',
    split='whitespace',
    output_mode='multi_hot')

train_text = raw_train_ds.map(lambda text,labels: text)
multi_hot_vectorize_layer.adapt(train_text)
```

The following Python code block illustrates the type of output generated by the text vectorization layer for the first element of the first batch of the training data set. The `next()` function gets the next element of an iterator, in this case an iterator over the training data set and the first element is a batch of text input and corresponding labels.

```
# Retrieve a batch from the dataset
text_batch, label_batch = next(iter(raw_train_ds))

# Applying the text vectorization layer
# to the first example and print its output
print(text_batch[0])
print(list(multi_hot_vectorize_layer(text_batch[0]).numpy()))
```

The first element of the `text_batch` is a tensor containing the raw text, while the output is a set of 10,000 numbers that represent the count of individual words (after standardization and tokenization).

```
>>> print(text_batch[0])
tf.Tensor(b'"unit testing of setters and getters teacher wanted us
to do a comprehensive unit test. for me, this will be the first time
that i use junit. i am confused about testing set and get methods.
do you think should i test them? if the answer is yes; is this code
enough for testing?... public void testsetandget(){.    int a = 10;.
class firstclass = new class();.    firstclass.setvalue(10);.    int
value = firstclass.getvalue();.    assert.asserttrue(""+error"",
value==a);. }...in my code, i think if there is an error, we can\'t
know that the error is deriving because of setter or getter.\n',
shape=(), dtype=string)

>>> print(list(multi_hot_vectorize_layer(text_batch[0]).numpy()))
[1.0, 1.0, 1.0, 1.0, 1.0, 1.0, 1.0, 1.0, 0.0, 1.0, 1.0,
1.0, 1.0, 0.0, 0.0, 0.0, 0.0, 0.0, 1.0, 1.0, 0.0, 0.0, 0.0,
0.0, 1.0, 1.0, 0.0, 1.0, 1.0, 1.0, 0.0, 0.0, 0.0, 0.0, 0.0,
1.0, 1.0, 1.0, 0.0, 0.0, 1.0, 0.0, 0.0, 0.0, 0.0, 1.0, 0.0,
0.0, 0.0, 0.0, 0.0, 1.0, 0.0, 0.0, 0.0, 0.0, 0.0, 1.0
...]
```

This simple example uses a classification model defined with a single fully-connected output layer. The four output units represent class probability logits. Cross-entropy is specified for the loss function, and accuracy metrics will be examined. Note the use of the `from_logits` parameter when specifying the loss function. This avoids the use of softmax activation function for the dense layer or adding a softmax layer to the sequential network model.

```
# Define a simple model
multi_hot_model = tf.keras.Sequential([
    multi_hot_vectorize_layer,
    layers.Dense(4)
])
# Set loss function, optimizer and metrics
multi_hot_model.compile(
    loss=losses.SparseCategoricalCrossentropy(from_logits=True),
    optimizer='adam', metrics=['accuracy'])
```

The training accuracy of this model is $\approx 99\%$ and the validation accuracy is $\approx 83\%$ showing the power of a relatively simple network architecture, albeit with significant number of trainable parameters (with 10,000 inputs from the vectorization pre-processing layer and four outputs, the network has 400004 trainable parameters).

Hands-On Exercise

Adapt the network architecture to identify the impact on training and validation performance of the following:

1. Vocabulary size (originally 10 000)
2. Number of layers (originally 1)

Comment on your findings and identify the best model.

14.7.2 Text classification using word embedding with a CNN

The example in this section illustrates a different way to encode words that is more space efficient while also accounting for the word sequence in a text. Instead of counting the number of times a word appears in the input, each word is first represented by a unique integer number.

The Keras `TextVectorization` layer can perform this, with a small change from the above example: Instead of the `multi_hot` output mode, the following code block specifies the `int` output mode for the layer:

```
# Use the Keras TextVectorization pre-processing layer:
int_vectorize_layer = TextVectorization(
    max_tokens=10000,
    standardize='lower_and_strip_punctuation',
    split='whitespace',
    output_mode='int')

train_text = raw_train_ds.map(lambda text, labels: text)
int_vectorize_layer.adapt(train_text)
```

The following Python code block illustrates the type of output generated by the text vectorization layer for the first element of the first batch of the training data set.

```
# Retrieve a batch from the dataset
text_batch, label_batch = next(iter(raw_train_ds))

# Applying the text vectorization layer
# to the first example and print its output
print(text_batch[0])
print(list(int_vectorize_layer(text_batch[0]).numpy()))
```

The printed output below illustrates how raw input text, the first element of the first batch of the training data set, is converted into a set of integer numbers by this layer.

The output is a list of 21 tokens (among the 10,000 most frequent ones). For example, the word "how" is coded as 5, the word "to" is coded as 4, etc.

```
>>> print(text_batch[0])
tf.Tensor(b'"how to crop a portion from an existing pdf and create
a new pdf in blank? i want to crop a portion in sizes like a5, a6
from an existing pdf of size a4 and want to create a new pdf"\n',
shape=(), dtype=string)

>>> print(list(int_vectorize_layer(text_batch[0]).numpy()))
[24, 4, 6757, 5, 2173, 31, 32, 1183, 962, 8, 124, 5, 15, 962, 7, 16,
3, 46, 4, 6757, 5, 2173, 7, 2661, 48, 6931, 1, 31, 32, 1183, 962, 9,
319, 2877, 8, 46, 4, 124, 5, 15, 962]
```

Word Embeddings However, these word numbers should not be treated as numerical data; they are categorical with no implicit ordering. Each word number is entirely arbitrary and the numbers could be permuted in any way. So rather than operate on token numbers this example uses *word embeddings* to generate an *embedding vector* of numbers of length k that maps each word to a point in a k dimensional space. That is, the set of numbers in the embedding vector characterizes or describes each word. The dimensionality of the space k can be chosen arbitrarily. Larger k can provide a better separation of many different unique words in the vector space, but require more computation, more memory, and may be more prone to overfitting. Smaller k may provide a poorer separation of words in the vector space, that is, multiple words are close together and difficult to differentiate. Essentially, word embeddings function as a *lookup table* where each row represents one word and the data in that row forms the embedding vector. Figure 14.17 shows a simple example of a word embedding table with $k = 4$ dimensions, that is, each word is represented by an embedding vector of length 4. In practice, dimensionality is often chosen as \log_{10} or a similar function of the input vocabulary size.

This example uses a sequential model beginning with the integer vectorization layer. An embedding layer with embedding size of 64 follows this. The input to the embedding layer is the output of the vectorization layer, that is, an integer for each token in the preprocessed input text. The embedding layer converts each word into a vector of 64 numeric values. The embedding layer is defined with a vocabulary size of 10,001 so that unknown tokens can be represented, that is, tokens not in the maximum 10,000 unique tokens of the vectorization layer. In effect, the embedding layer is a large lookup table with 10,001 rows and 64 columns. Hence, it contains 64,064 trainable parameters. The `mask_zero` parameter for the embedding layer instructs the layer to treat the number 0 as a padding value, rather than as the number for a specific word. Because the length of the word sequence can vary with the length of the input text, padding inputs with 0 ensures that all inputs in batches are of the same size.

A dropout layer with a dropout rate of 0.5 is added to the sequential model for regularization. This is followed by a 1-dimensional convolution layer with kernel length of 5. The Conv1D layer uses 64 separate filters to produce 64 output channels from an input

A 4-dimensional embedding

cat =>	1.2	-0.1	4.3	3.2
mat =>	0.4	2.5	-0.9	0.5
on =>	2.1	0.3	0.1	0.4

...

...

https://www.tensorflow.org/text/guide/word_embeddings

Figure 14.17: Example of word embedding

depth of 64. The global max pooling layer reduces the input to the maximum value along each convolution output channel.

```
int_model = tf.keras.Sequential([
    int_vectorize_layer,
    layers.Embedding(10001, 64, mask_zero=True),
    layers.Dropout(0.5),
    layers.Conv1D(filters=64, kernel_size=5, \
        padding="valid", activation="relu", strides=2),
    layers.GlobalMaxPooling1D(),
    layers.Dense(4)
])
int_model.summary()
```

It is instructive to examine the model summary, printed below. Note that the output shapes have no value for the first dimension, that is, the batch size. This will be determined when the model is trained using the batch size provided by the training data set (The above code segment when preparing the data set defined the batch size as 32). The second dimension of the output shapes for any layer that processes sequences of words is also not explicitly defined. This second dimension indicates the sequence length. It will be determined from at training time when the training data set is provided as input, potentially padded by 0 if necessary. Note that the global max pooling layer reduces the dimensionality as its output is the maximum value along the second dimension, that is, the maximum value over the sequence of convolutional outputs.

The model summary output below also confirms the number of parameters of the word embedding layer. From the parameter count of the 1-dimensional convolution layer, one can deduce the specific kernel size. Knowing there are 64 filters one can subtract one bias term for each filter, leaving 20,480 parameters for 64 output channels. That is, each output channel is calculated by a kernel with 320 parameters. This is because

these are one-dimensional kernels of length 5 and the input (output of the dropout layer) is of depth 64, that is, the full kernel must be of shape 5×64 with 320 parameters.

```
Model: "sequential_2"
-----  

Layer (type)          Output Shape         Param #
-----  

text_vectorization_1 (Text    (None, None)           0  

embedding (Embedding)     (None, None, 64)        640064  

dropout_2 (Dropout)       (None, None, 64)           0  

conv1d (Conv1D)          (None, None, 64)        20544  

global_max_pooling1d (Glob (None, 64)              0  

dense_6 (Dense)          (None, 4)                260  

-----  

Total params: 660868 (2.52 MB)  

Trainable params: 660868 (2.52 MB)  

Non-trainable params: 0 (0.00 Byte)
```

Next, the model is compiled with cross-entropy as loss function and accuracy as a result metric:

```
# Compile the model using cross-entropy loss and report accuracy
int_model.compile(
    loss= losses.SparseCategoricalCrossentropy(from_logits=True),
    optimizer='adam',
    metrics=['accuracy'])
```

Finally, the model is trained for 25 epochs:

```
int_model.fit(raw_train_ds,
              validation_data=raw_val_ds,
              epochs=25)
```

The training accuracy of this model is $\approx 99.95\%$ but with a validation accuracy similar to the earlier, simpler model of $\approx 80\%$, showing very good training performance but also more evidence of overfitting. To some degree this is unsurprising, as the number of parameters in this model is about 50 times that of the simpler model, making this model more powerful and more flexible to adapt to the specific training set. This short example illustrates another use of convolutional networks besides computer vision, where they were first used.

The Keras web site has a large number of Keras tutorials on all kinds of natural language processing problems. See this site for more information: <https://keras.io/examples/nlp/>.

Hands-On Exercise

Adapt the network architecture to identify the impact on training and validation performance of the following:

1. Vocabulary size (originally 10 000)
2. Embedding size (originally 64)
3. Dropout probability (originally 0.5)
4. Convolution kernel size (originally 5)
5. Number of convolution filters (originally 64)
6. Convolution stride (originally 2)
7. Number of 1D-Conv layers (originally 1)

Comment on your findings and identify the best model.

In particular, explore the trade-offs between using a large number of smaller filters versus a smaller number of larger filters in the context of convolution followed by pooling.

Explore how does the choice of pooling strategy (max vs average) affects the features extracted by a convolutional neural network and the predictive performance of the model.

14.8 Review Questions

Introduction to CNNs

1. What are Convolutional Neural Networks (CNNs) commonly used for in the field of computer vision?
2. Explain the primary motivation for using convolutional networks over fully connected networks in image processing.
3. Explain why convolutional layers are more efficient than fully connected layers in the context of image data.
4. Describe the typical structure of a CNN in terms of how it processes features at different levels of abstraction.
5. Illustrate with examples how CNNs can be used in automated surveillance systems.
6. Discuss the role of CNNs in enhancing user experiences in augmented reality applications.
7. Explain the challenges and considerations in using CNNs for medical image analysis, particularly in ensuring reliability and accuracy.
8. How do advancements in CNN technologies potentially impact the development of autonomous driving technologies?

Convolutional Layers

9. Define a convolutional layer. What role do the filters (kernels) play within such a layer?
10. Explain the convolution operation in the context of a CNN. How does it differ from the operations performed by traditional neural networks?
11. What is meant by an ‘activation map’ or ‘feature map’ in the context of CNNs?
12. Discuss the purpose of striding and padding in convolution operations, providing examples of their effects on the output dimensions.
13. Given a CNN with an input size of $32 \times 32 \times 3$ (where 3 stands for RGB channels) and a convolutional layer with 10 filters of size $5 \times 5 \times 3$, calculate the output dimensions assuming a stride of 1 and no padding. Also, discuss how the output dimensions would change with a stride of 2 and padding size of 1.
14. Describe the roles of biases and activation functions in the context of convolutional layers. What is a commonly used activation function in CNNs?
15. Refer to Figure 14.1 and explain the process depicted in a 1-dimensional convolution example.
16. Analyze Figure 14.2 to describe how 2-dimensional convolution differs from 1-dimensional convolution.
17. Explain how padding can influence the learning process in CNNs, particularly in relation to border information in images.
18. Discuss the potential benefits and drawbacks of using large strides in convolutional layers.
19. What are the advantages of using small kernel sizes (e.g., 3×3) in CNN layers, and how might using such kernels affect the learning capacity of the network?

Pooling

20. Define pooling in the context of Convolutional Neural Networks. What is its main purpose?
21. Compare and contrast max pooling and average pooling. How does each affect the feature map it processes?
22. Explain how pooling layers contribute to the invariance of ConvNets to small transformations, distortions, and translations in the input.
23. Discuss the impact of pooling size and stride on the dimensionality of the output. How do these parameters influence the model’s ability to detect fine-grained features versus more abstract features?
24. Discuss the advantages and potential drawbacks of using overlapping pooling regions in ConvNets.
25. Can pooling layers lead to loss of important information? Provide reasons for your answer and discuss how this might affect the overall performance of a CNN.
26. Explain why pooling might be useful in terms of computational efficiency and model generalization.

Object Detection

27. Explain how CNNs can be applied to the task of object detection. Include a discussion of how bounding or anchor boxes are used.

28. Compare and contrast the approaches of R-CNN and YOLO in object detection. How do they handle the detection task differently?
29. Discuss the significance of anchor boxes in modern object detection models like YOLO and SSD.

Semantic Segmentation

30. How does semantic segmentation differ from object detection in terms of output and application areas?
31. How does integrating object detection with semantic segmentation enhance the performance of a vision system in autonomous driving scenarios?
32. Evaluate the challenges in training CNNs for semantic segmentation of irregular objects, such as trees or clouds. What techniques can be used to address these challenges?
33. Evaluate the impact of image resolution on the performance of semantic segmentation. How does it affect the accuracy of pixel classification?
34. How can real-time constraints influence the design of CNN architectures for object detection and semantic segmentation in systems like autonomous vehicles?
35. Discuss the computational trade-offs involved in running both object detection and semantic segmentation in real-time applications such as video surveillance.
36. Consider the ethical implications of deploying CNN-based object detection and semantic segmentation technologies in public spaces. What privacy concerns arise, and how can they be mitigated?

Word Embeddings

37. Discuss the advantages of using word embeddings over traditional one-hot encoding in natural language processing.
38. How does the dimensionality k of word embeddings affect the performance of neural networks in NLP tasks? Include a discussion of the trade-offs involved with choosing a higher or lower k .
39. Describe how an embedding layer works in the context of a neural network. What are the trainable parameters within an embedding layer?
40. Describe the role of padding in text sequences when using embedding layers. How does this affect the processing of batched text data?
41. Explain the interaction between the embedding layer and subsequent layers in a neural network, such as a 1-dimensional convolutional layer. How does the output of the embedding layer serve as input to the convolutional layer?
42. How can embeddings be shared across different tasks or models in machine learning? Discuss the advantages and potential issues with this approach.
43. Consider the ethical implications of using word embeddings in text processing. What biases might be inherent in pretrained embeddings, and how can they affect the outcomes of NLP applications?

Chapter 15

Recurrent Neural Networks

Sources and Further Reading

Kevin P. Murphy: *Probabilistic Machine Learning – An Introduction*. MIT Press 2022.

<https://probml.github.io/pml-book/book1.html>

Chapter 15

The book by Murphy is freely available and provides three chapters on neural networks, one for structured data, one for images, and one for sequences. It provides significant depth on convolutional and recurrent network architectures, fitting the models, and problems the data analyst may encounter.

Guides and examples on the Tensorflow and Keras web sites:

- [Time Series Forecasting \(Tensorflow\)](#)
- [Working with RNNs \(Tensorflow\)](#)
- [Text Generation with an RNN \(Tensorflow\)](#)
- [Timeseries Forecasting for Weather Prediction \(Keras\)](#)

This course uses the Tensorflow programming framework for neural network applications. The Tensorflow website has a multitude of introductory and advanced guides and tutorial that cover all aspects of machine learning with neural networks.

Introductory tutorials:

Olah, Christopher (2015) [Understanding LSTM Networks](#)

Karpathy, Andrej (2015) [The Unreasonable Effectiveness of Recurrent Neural Networks](#)

Chris Olah has been lead researcher at OpenAI and Google Brain and co-founded Anthropic. Andrej Karpathy has been lead researchers at OpenAI and Tesla. Both tutorials are very useful and easy introductions to the topic of recurrent LSTM networks.

15.1 Introduction

Recurrent Neural Networks (RNNs) are a class of neural networks that are used for modeling sequence data such as time series, natural language, or audio. Characterized by their ability to maintain a "memory" of previous inputs while processing new ones, RNNs are particularly useful for tasks where historical context is important.

The development of RNNs can be traced back to the 1980s with the introduction of architectures that could use their internal state (memory) to process sequences of inputs. This concept was refined in the 1990s through the introduction of the Long Short-Term Memory (LSTM) network, which significantly improved the performance of RNNs on tasks requiring learning long-term dependencies.

RNNs are employed in a variety of applications, handling different types of data and tasks:

- *Natural Language Processing (NLP)*: RNNs are the basis for many NLP tasks such as machine translation, speech recognition, and text generation. Their ability to process sequences of words and maintain context helps in producing better translations and recognizing speech accurately.
- *Time Series Prediction*: RNNs are suitable for forecasting future events in financial markets, weather conditions, and other time-dependent phenomena due to their ability to remember past data.
- *Music and Video Generation*: By learning from sequences of musical notes or frames of videos, RNNs can generate new music pieces and video clips that are similar in style to their training data.
- *Audio Transcription and Video Captioning*: By examining sequences of audio or visual images, RNNs can generate text transcripts or captions of the video content.
- *Healthcare*: In medical diagnostics, RNNs can predict disease progression and patient outcomes by analyzing sequential data such as patient records and time-series observations from medical sensors.

- *Business Processes:* In business processes, RNNs can be used to predict the next activity, potential problems, time to completion, or other outcomes based on the series of actions already completed.

15.2 Sequence Models

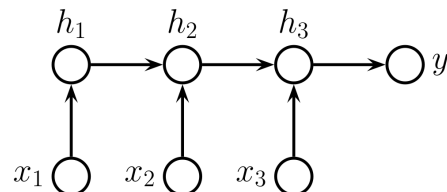
Recurrent Neural Networks (RNNs) are called "recurrent" because they perform the same task for every element of a sequence, with the output being dependent on the previous computations. RNNs can pass information from one (time or sequence) step of the network to the next. This mechanism is what makes RNNs "recurrent" – they recur or repeat the same process over each part of the input, while maintaining some memory of what has happened before. This memory is maintained through internal or "hidden" states of the network, which capture information about earlier elements in the sequence, thereby providing a form of memory. This allows RNNs to process not just individual data points, but entire sequences of data (such as a sentence or a time series), making them effective for tasks where context and order are important.

In using RNNs with sequence data, one can distinguish three types of general model architectures that are useful for different tasks.

Seq2Vec

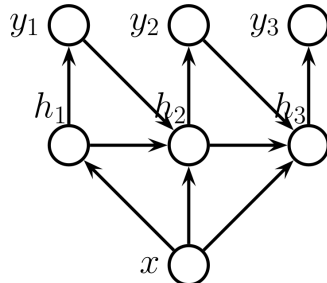
The sequence-to-vector ("Seq2Vec") model predicts a single outcome or target from a sequence of inputs. This type of model can be a regression or classification model. The general model architecture is shown in Figure 15.1. The hidden states h_i are connected to model the memory through time. The state of the network depends not only on the current input but also on the previous state, and therefore implicitly on all prior inputs. In practical applications, the inputs, outputs, and hidden state can be high-dimensional vectors or arrays.

This type of architecture is useful for example in sentiment analysis, where it can analyze sequences of text to determine the sentiment expressed, summarizing the overall sentiment in a single vector for classification.



Source: Murphy Fig. 15.4

Figure 15.1: Seq2Vec recurrent neural network architecture



Source: Murphy Fig. 15.1

Figure 15.2: Vec2Seq recurrent neural network architecture

Vec2Seq

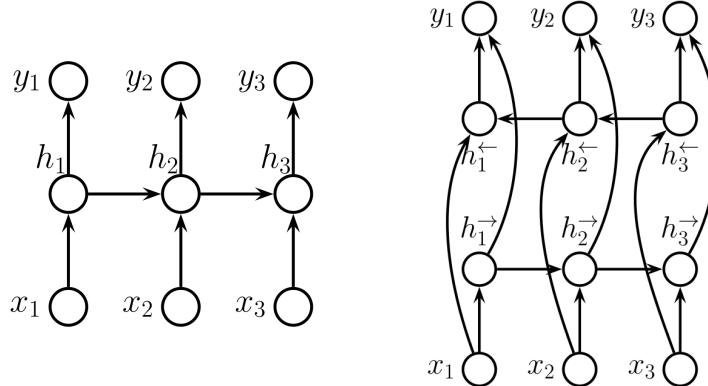
The vector-to-sequence (“*Vec2Seq*”) models are designed to generate a sequence from a fixed-length input vector. This approach is commonly used in tasks where a sequence needs to be generated from a compact representation. Figure 15.2 shows an example of such an architecture. A single input x gives rise to multiple outputs y with the hidden states h maintaining information about the history. Here, the state of the network depends not only on the single input but also on the previous state and previous output.

The Vec2Seq architecture is useful for example in image captioning, generating a sequence of words as descriptive text for a single input image (a vector of pixels). Another use case is music generation, creating a sequence of musical notes from an input vector that encodes a particular style or mood.

Seq2Seq

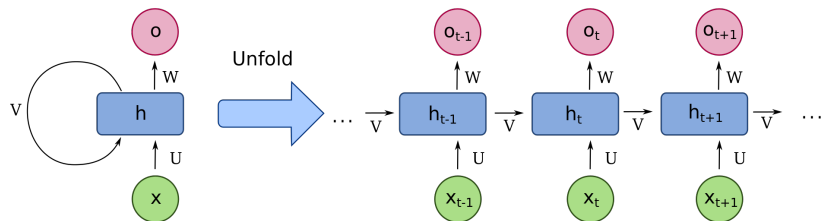
Sequence-to-sequence (“*Seq2seq*”) models are designed to transform an input sequence into an output sequence. These models are useful for tasks that involve translation or conversion from one sequence to another, maintaining the context from the input to the output. The general architecture is shown in Figure 15.3. For each input x , one output y is generated and the hidden state h maintains information about the history. The state of the network depends not only on the current input, but also on the previous state. Depending on the application, Seq2Seq networks may also employ *bi-directional connections* between hidden layers, as shown in the right panel of Figure 15.3.

A typical use case for Seq2Seq models is machine translation from one language to another, where both the input and output are sequences of words. In speech recognition, spoken language is converted into written text, where the audio input is a sequence of phonetic features, and the output is a sequence of words. In video-to-text applications, textual descriptions are generated from video sequences, which involves interpreting sequences of images and producing corresponding sequences of descriptive text.



Source: Murphy Fig. 15.5

Figure 15.3: Seq2Seq recurrent neural network architecture



https://commons.wikimedia.org/wiki/File:Recurrent_neural_network_unfold.svg

Figure 15.4: Unrolling a recurrent neural network

15.3 Unrolling an RNN

The concept of "unfolding" or "unrolling" in recurrent neural networks (RNNs) is a fundamental technique used not only to visualize but more importantly, to implement these networks for sequence processing. Unrolling an RNN refers to the process of expanding the recurrent network through time, transforming it into an equivalent feed-forward neural network that represents each time or sequence step explicitly. This helps in understanding and analyzing the behavior of RNNs, especially in training.

An RNN is designed to handle sequences by recursively processing each element of the sequence, maintaining a hidden state that captures information about the past elements of the sequence. When RNN is unrolled, each recurrence becomes a separate, but identical, unit of the network. Each unit corresponds to a time step in the input sequence.

Consider the example in Figure 15.4. The recursive network on the left feeds back

information to itself by vector v . This information updates the hidden layer's state h from one recurrence (time or sequence step) to the next. This recursive network is unrolled to separate instances of input, hidden layers, and output for each sequence step or time step as shown in the right part of Figure 15.4. The hidden state h and output o in Figure 15.4 can be described using basic neural network units:

$$h_t = \sigma(W_x \cdot x_t + W_h \cdot h_{t-1} + B_h) \quad (15.1)$$

$$o_t = \sigma(W_o \cdot h_t + B_o) \quad (15.2)$$

For example, if the input are sequences of five words and an RNN is designed to process this sequence, unrolling this RNN would result in a chain of five identical neural network units (one for each word). Each unit takes as input the current word, however encoded, and the hidden state output by the previous unit. The first block receives an initial hidden state, often set to zero, which is a starting state.

Unrolling makes the sequence processing capabilities of RNNs explicit and easier to understand, showing how inputs are processed over time. In practice, unrolling an RNN simplifies its implementation, especially for training where gradients are computed across the unfolded network.

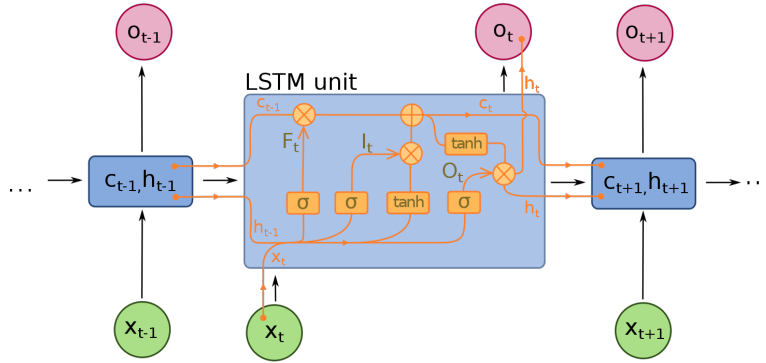
Unrolling an RNN over many time steps can lead to practical challenges. In particular, long unrolled networks with simple units as in Equations 15.1 and 15.2 often suffer from vanishing or exploding gradient problems during training, which makes learning unstable. The vanishing gradient problem in particular means that the network loses its "memory" of inputs in the distant past. In other words, such a network is unable to have "long term" memory that is useful for many applications.

15.4 LSTM Cells

Long Short-Term Memory (LSTM) cells are a special kind of unit used in recurrent neural networks (RNNs) that are designed to address some of the limitations of traditional RNNs. In particular, they enable networks to learn long-term dependencies that are critical in many sequential tasks such as language modeling or time series prediction. LSTMs were introduced in 1997 as a solution to the vanishing gradient problem commonly encountered in traditional RNNs. Traditional RNNs could perform well on short sequences but struggled with longer ones.

An LSTM cell, shown schematically in Figure 15.5, is complex and contains multiple "gates" that regulate the flow of information. LSTM networks have two components to their state. The *cell memory* is represented by c_t and the hidden state is represented by h_t in Figure 15.5. Each LSTM cell consists of the following components:

- *Forget Gate*: This gate decides which information is discarded from the cell memory c . It looks at the previous hidden state and the current input and passes its output through a sigmoid function, which outputs numbers between 0 ("forget



https://en.wikipedia.org/wiki/File:Long_Short-Term_Memory.svg

Figure 15.5: Long Short-Term Memory Cell

this vector element completely") and 1 ("keep this vector element entirely"). The forget gate is represented by F_t in Figure 15.5 and formally defined as:

$$F_t = \sigma(W_f \cdot [x_t, h_{t-1}] + b_f) \quad (15.3)$$

- *Input Gate:* This gate updates the cell memory c by adding new information. It looks at the previous hidden state and the current input and includes a sigmoid layer which decides which values to update. It outputs numbers between 0 ("do not update this vector element") and 1 ("update this vector element completely"). This input gate is represented by I_t in Figure 15.5 and formally defined as:

$$I_t = \sigma(W_i \cdot [x_t, h_{t-1}] + b_i) \quad (15.4)$$

- *Output Gate:* The output gate examines the prior hidden state and current input. It includes a sigmoid layer which decides which values of the new cell memory to output and pass on to the next cell as the new hidden state. It outputs numbers between 0 ("do not include this vector element in the output") and 1 ("include this vector element in the output"). The output gate is shown as O_t in Figure 15.5 and formally defined as:

$$O_t = \sigma(W_o \cdot [x_t, h_{t-1}] + b_o) \quad (15.5)$$

The outputs of these three gates are used to manipulate the cell memory c_{t-1} that was received from the previous LSTM unit. First, a tanh unit creates a new candidate vector cell memory vector (not explicitly labeled in Figure 15.5), formally defined as:

$$\tilde{c}_t = \phi(W_c \cdot [x_t, h_{t-1}] + b_c) \quad (15.6)$$

The new cell memory c_t is created by first multiplying the old cell memory c_{t-1} with the output of the forget gate O_t (Eq. 15.3), thus removing some information. Second,

information from the candidate cell memory (Eq. 15.6) is selected by multiplying it with the values of the input gate I_t (Eq. 15.4) and is then added to the new cell memory. Formally:

$$c_t = F_t \otimes c_{t-1} + I_t \otimes \tilde{c}_t \quad (15.7)$$

The output of the LSTM cell is also the new cell state h_t . It is formed by applying the output gate O_t (Eq. 15.5) to the cell memory. That is, it selectively outputs and passes on parts of the new cell memory as determined by the output gate. It is formally defined as:

$$h_t = O_t \otimes \phi(c_t) \quad (15.8)$$

In Equations 15.3 to 15.8, \cdot denotes the dot-product (vector product), \otimes denotes element-wise multiplication, and $[.]$ denotes vector concatenation. σ is the sigmoid/logistic function and ϕ is the hyperbolic tangent (\tanh).

15.5 GRU Cells

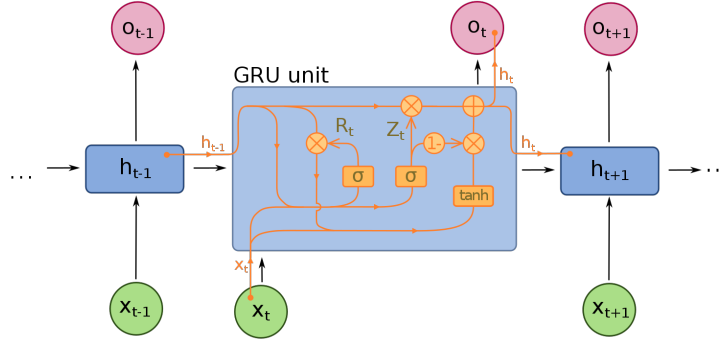
Gated Recurrent Unit (GRU) cells are a type of recurrent neural network (RNN) architecture introduced as an alternative to Long Short-Term Memory (LSTM) cells. The motivation behind the development of GRUs was to simplify the LSTM architecture, which, although powerful, is also quite complex due to its multiple gates and states. By reducing the number of gates from three to two, GRUs aim to offer a model that can train faster and require fewer computational resources, while still capturing long-range dependencies within the data. GRUs also do not have an internal memory unit separate from the hidden state. While GRUs tend to train faster due to their simpler structure, in some cases LSTMs outperform GRUs in predictive performance, if the additional complexity of LSTMs are appropriate to the data set and task.

A GRU cell, schematically shown in Figure 15.6 consists of only two gates, the *reset gate* and the *update gate*. It also merges the cell memory and the hidden state.

- *Reset Gate*: This gate, represented as R_t in Figure 15.6, determines how much past information to forget, which is important for making the model more adaptable to changes in the data sequence. It examines the previous state and current input and applies a sigmoid function. It outputs values between 0 ("reset this vector element completely") and 1 ("retain this vector element completely").

$$R_t = \sigma(W_r \cdot [x_t, h_{t-1}] + b_r) \quad (15.9)$$

- *Update Gate*: This gate, represented as Z_t in Figure 15.6, decides how much of the past information (from previous time or sequence steps) needs to be passed



https://en.wikipedia.org/wiki/File:Gated_Recurrent_Unit.svg

Figure 15.6: Gated Recurrent Unit (GRU)

along to the future. It is similar to the combination of the forget and input gates in an LSTM. It examines the previous state and current input, applies a sigmoid function and outputs values between 0 ("do not update this vector element, retain the old state") and 1 ("update this vector element, discard the old state").

$$Z_t = \sigma(W_z \cdot [x_t, h_{t-1}] + b_z) \quad (15.10)$$

The candidate new hidden state is calculated by applying the reset gate to the prior hidden state and then combining the result with the current input and applying a tanh activation function. This "forgets" certain information from the prior state. Formally:

$$\hat{h}_t = \phi(W_h \cdot [x_t, R_t \otimes h_{t-1}] + b_h) \quad (15.11)$$

Finally, the actual new hidden state is calculated as a weighted sum of the past hidden state, weighted by $1 - Z_t$, and the candidate state, weighted by Z_t .

$$h_t = (1 - Z_t) \otimes h_{t-1} + Z_t \otimes \hat{h}_t \quad (15.12)$$

In Equations 15.9 to 15.12, \cdot denotes the dot-product (vector product), \otimes denotes element-wise multiplication, and $[.]$ denotes vector concatenation. σ is the sigmoid/logistic function and ϕ is the hyperbolic tangent.

15.6 Statefulness

Recurrent Neural Networks (RNNs) are designed to handle sequence data. One of the key decisions in designing RNNs is whether to use a stateful or stateless configuration. This choice affects how the network processes sequences and learns from data, influencing both the training process and application effectiveness.

In *stateless RNNs*, the internal state is reset for each batch of training data. This means that the learning and predictions are independent of the previous batch. The network starts with a clean slate for every input sequence, making no assumptions based on past data points within the same batch. Since stateless RNNs treat each input sequence as independent, training can be simpler and faster. There is no need to maintain and manage state continuity across batches, and batches can be assembled from the input sequences in arbitrary order and randomly shuffled. Stateless RNNs are typically used when sequences are short enough to fit within the number of unrolled RNN steps and the context within a single sequence is sufficient for the prediction task.

Stateful RNNs maintain the internal state between batches, allowing the network to retain information across different input batches. This is useful when sequences are longer than the number of unrolled steps. Training stateful RNNs requires careful management of the state so that it is carried over appropriately across batches. Batches cannot be randomly drawn and cannot be shuffled. The state must be reset manually at the start of each epoch or when starting a new sequence. Stateful RNNs are advantageous for tasks involving long sequences where the context needs to be maintained over long periods, such as in time-series analysis or long-form text processing.

15.7 Example – Stock Market Prediction

This example uses historic stock market data to predict future performance. In particular, it uses data for the Dow Jones Industrial Average (DJIA) that was exported from the `quarks` library for R. The data set encompasses opening closing, low, and high values, as well as trading volume for every day between the years 2000 and 2021 (converted to Euros). This is a regression problem with the targets being the difference in closing prices to the next day.

Complete implementations for this and the other examples in this chapter are available in the following GitHub repo: <https://github.com/jevermann/busi4720-ml>

The project can be cloned from this URL: <https://github.com/jevermann/busi4720-ml.git>

First, load all required packages, specify parameters for the network architecture and training, and then read the data file:

```

import math
import tensorflow as tf
from tensorflow import keras
from keras import layers
import pandas as pd

tf.random.set_seed(123)      # Set random seed
n_steps = 20                  # Unfold for 20 steps
n_epochs = 25                  # Train for 25 epochs

# The read the data set
data = pd.read_csv('https://evermann.ca/busi4720/djia.data.csv')

```

In the next step, additional features are created. In time series analysis it is often useful to examine the differences between successive values and the relative or percentage changes. The following Python code fragment adds these to the data set by applying the Pandas functions `diff()` and `pct_change()` to the columns of the data frame and concatenating the differences and percentage changes to the dataframe by column.

```

data = pd.concat([data,
    data.diff().add_suffix('diff'),
    data.pct_change().add_suffix('pct')], 
    axis=1).iloc[1:,]

```

When splitting the data into training and validation set, random shuffling cannot be used because the data set is time series data. In other words, information from future observations should not be used in the training set to help predict values of past observations. The following Python code uses the first 80% of the data set for training and the remainder for validation:

```

# Split data, no random shuffling for time series
train = data[:math.floor(0.8*data.shape[0])]
valid = data.drop(train.index)

```

Normalizing the data to zero means and unit standard deviation must again only use information from the training data set, even if that means the actual validation set standard deviations are not exactly equal to 1:

```

# Normalize data using only info from training set.
train_mean = train.mean()
train_sd = train.std()
train = (train - train_mean)/train_sd
valid = (valid - train_mean)/train_sd

```

An easy way to create appropriate sequence batches and targets is to use the Keras function `timeseries_dataset_from_array()`. It accepts the features, the

targets`m` and arguments about the sequence length, batch size, and whether the data should be shuffled between batches. This first example will use a stateless LSTM so that the input can be shuffled. This is a regression problem with the target being the closing price difference to the following day.

```
dataset_train = keras.preprocessing.timeseries_dataset_from_array(
    train.drop('price.closediff', axis=1), train['price.closediff'],
    sequence_length=n_steps,
    batch_size=32,
    shuffle=True)

dataset_valid = keras.preprocessing.timeseries_dataset_from_array(
    valid.drop('price.closediff', axis=1), valid['price.closediff'],
    sequence_length=n_steps,
    batch_size=32,
    shuffle=True)
```

Tensorflow Datasets for Sequence Data

To see how Tensorflow manages time series or sequence data in its datasets, it is instructive to work with a simple example. Consider the following Pandas dataframe of inputs and targets:

```
test = pd.DataFrame([[0, 0], [1, 1], [2, 2], [3, 3],
                     [4, 4], [5, 5], [6, 6], [7, 7]])
inputs = test.iloc[:-1,0]
targets = test.iloc[2:,1]

pd.DataFrame([inputs, targets])
```

	0	1	2	3	4	5	6	7
0	0.0	1.0	2.0	3.0	4.0	5.0	6.0	NaN
1	NaN	NaN	2.0	3.0	4.0	5.0	6.0	7.0

The first example illustrates creating a dataset object that provides batches of size 2 for a sequence length of 2, using a stride of 1 across the input sequences. This example does not shuffle the data between batches and is therefore suitable for stateful RNNs:

```
ds = keras.preprocessing.timeseries_dataset_from_array(
    inputs, targets,
    batch_size=2, sequence_length=2, sequence_stride=1, shuffle=False)
for element in ds.as_numpy_iterator():
    print(element)
```

The following output shows batches of sequences and their targets. Note how the sequences are continuous across batches. For example the first sequence of 2 elements

of the first batch is [0, 1] and the first sequence of 2 elements of the second batch is [2, 3], that is, the second batch continues sequences from the first batch, and the hidden state in the LSTM or GRU cells in the RNN can be carried over to the next batch.

```
(array([[0, 1],
       [1, 2]]), array([2, 3]))
(array([[2, 3],
       [3, 4]]), array([4, 5]))
(array([[4, 5],
       [5, 6]]), array([6, 7]))
```

In contrast, the next example shuffles elements between batches and is suitable for stateless RNNs where state is not maintained between batches:

```
ds = keras.preprocessing.timeseries_dataset_from_array(
    inputs, targets,
    batch_size=2, sequence_length=2, sequence_stride=1, shuffle=True)
for element in ds.as_numpy_iterator():
    print(element)
```

The output below clearly shows the differences. The sequences of length 2 are not continuous across batches. For example, the first sequence of 2 elements in the second batch is [5, 6] while the first sequence of 2 elements in the third batch is [3, 4].

```
(array([[4, 5],
       [2, 3]]), array([6, 4]))
(array([[5, 6],
       [0, 1]]), array([7, 2]))
(array([[3, 4],
       [1, 2]]), array([5, 3]))
```

The next example shows the effect of varying the sequence stride and the batch size. The batch size is now set to 1, and the sequence stride is 2, that is, after assembling a series of 2 elements for a batch, the next set of 2 elements is taken from two positions further in the input.

```
ds = keras.preprocessing.timeseries_dataset_from_array(
    inputs, targets,
    batch_size=1, sequence_length=2, sequence_stride=2, shuffle=False)
for element in ds.as_numpy_iterator():
    print(element)
```

```
(array([[0, 1]]), array([2]))
(array([[2, 3]]), array([4]))
(array([[4, 5]]), array([6]))
```

In this small illustrative example, every step of the feature sequence is a single number, whereas in the stock market prediction example every step of the feature sequence is itself a vector of values.

Hands-On Exercise

Using the simple example data from above to:

1. Experiment with different values for `batch_size`,
2. Experiment with different values for `sequence_length`,
3. Experiment with different values for `sequence_stride`.

The stock market prediction example uses a sequential neural network model with an input layer to specify the shape of the features to be used for training, one LSTM layer and a dense (fully-connected) layer with a single output for regression. It has the following characteristics:

- Hidden state and cell memory size: `units=16`
- "Seq2Vec" model: `return_sequences=False`
- Stateless model: `stateful=False`

Note that the second dimension of the input shape is one less than the total number of columns in the training data, because the column of target values has been removed.

```
model = keras.Sequential()
model.add(layers.InputLayer(
    input_shape=(n_steps, len(train.columns)-1)))
model.add(layers.LSTM(
    units=16,
    return_sequences=False,
    return_state=False,
    stateful=False))
model.add(layers.Dense(1))
model.summary()
```

The model summary is shown below. It is instructive to verify the number of parameters. Weights and biases are shared between time steps; recall that the RNN actually has a single LSTM cell and unrolling is simply there to make the computation easier. That is, the total parameter count reported below is for a single LSTM cell. Equations 15.3 to 15.6 show that the forget, input and output gates as well as the candidate new memory of the LSTM unit use the same number of parameters, that is, each equation introduces $1984/4 = 496$ parameters. The Equations 15.3 to 15.6 operate on the concatenation of the hidden state and the input, that is on a vector of $16 + 14 = 30$ units. Each equation adds a bias term. Hence, the total number of weights and bias for each gate as $(16 + 14 + 1) * 16 = 496$.

```
Model: "sequential_4"

Layer (type)          Output Shape         Param #
=====
lstm_1 (LSTM)        (None, 16)           1984
dense_8 (Dense)      (None, 1)            17
=====
Total params: 2001 (7.82 KB)
Trainable params: 2001 (7.82 KB)
Non-trainable params: 0 (0.00 Byte)
```

Compile and fit the model:

```
model.compile(loss='mean_squared_error', optimizer='Adagrad')

model.fit(dataset_train, epochs=n_epochs,
          validation_data=dataset_valid)
```

The training output, shown below, indicates that the training loss after 25 epochs is ≈ 1.00 and the validation is ≈ 7.06 . Importantly, neither training nor validation loss have decreased during training, suggesting that either the neural network model is inappropriate or the problem of stock market prediction is quite difficult.

```
Epoch 1/25
138/138 [=====] - 2s 6ms/step -
loss: 1.0933 - val_loss: 7.0730

Epoch 25/25
138/138 [=====] - 1s 5ms/step -
loss: 1.0028 - val_loss: 7.0621
```

Hands-On Exercise

Download the Python code from [here](#). Then:

1. Predict the percentage change of the closing value (column `price.closepct`)
2. Predict the actual closing value (column `price.close`)
3. Comment on the model performance results. Are these values more or less predictable than the differenced closing values?

Experiment with different model characteristics:

1. Vary the size of the LSTM hidden state and output (`unit=16`).
2. Swap the LSTM layer for a GRU layer (`layers.GRU`). The Keras GRU layer construction function takes the same arguments as the Keras LSTM layer construction function.
3. Comment on the model performance results.

Given the poor performance of the single layer LSTM network, it is natural to ask whether more LSTM layers can make a difference. To illustrate the use of Seq2Seq models, the following model stacks two LSTM layers. Note that the first LSTM layer is a Seq2Seq type that returns an output for each step of the sequence. These outputs form the inputs for the second LSTM layer. That second layer is a Seq2Vec model that returns a single output. The model also adds a second fully-connected layer and dropout layers to prevent overfitting.

```
model = keras.Sequential()
model.add(layers.InputLayer(
    input_shape=(n_steps, len(train.columns)-1)))
model.add(layers.LSTM(
    units=16, return_sequences=True, stateful=False))
model.add(layers.Dropout(rate=0.25))
model.add(layers.LSTM(
    units=16, return_sequences=False, stateful=False))
model.add(layers.Dense(32))
model.add(layers.Dropout(rate=0.25))
model.add(layers.Dense(1))
model.summary()
```

The model summary and the final epoch training output are shown below. Unfortunately, the model performs no better than the simple one.

Layer (type)	Output Shape	Param #
lstm_2 (LSTM)	(None, 20, 16)	1984
dropout_3 (Dropout)	(None, 20, 16)	0
lstm_3 (LSTM)	(None, 16)	2112
dense_9 (Dense)	(None, 32)	544
dropout_4 (Dropout)	(None, 32)	0
dense_10 (Dense)	(None, 1)	33

Total params: 4673 (18.25 KB)
Trainable params: 4673 (18.25 KB)
Non-trainable params: 0 (0.00 Byte)

Epoch 25/25
138/138 [=====] - 2s 17ms/step
- loss: 1.0064 - val_loss: 7.1526

Perhaps more history than the last 20 steps is required for good predictions. The following example illustrates a stateful LSTM. First, change the batch size to 1 for a single sequence of continuous elements and prohibit shuffling of the data set:

```
dataset_train = keras.preprocessing \
    .timeseries_dataset_from_array(
        train.drop('price.closediff', axis=1),
        train[['price.closediff']],
        sequence_length=n_steps,
        sampling_rate=1, batch_size=1, shuffle=False)

dataset_valid = keras.preprocessing \
    .timeseries_dataset_from_array(
        valid.drop('price.closediff', axis=1),
        valid[['price.closediff']],
        sequence_length=n_steps,
        sampling_rate=1, batch_size=1, shuffle=False)
```

The neural network architecture is a single LSTM as in the initial example, but this time the LSTM layer is stateful. Note that the input layer does not specify a sequence length for the batch input shape.

```
model = keras.Sequential()
model.add(layers.InputLayer(
    batch_input_shape=(1, None, len(train.columns)-1)))
model.add(layers.LSTM(units=16,
    return_sequences=False, return_state=False, stateful=True))
model.add(layers.Dense(1))
model.summary()
```

Again, this model does not perform any better than the previous two models. As a baseline comparison, a Vec2Vec model is fitted with a single large feature vector that contains all 20 sequence steps. A Flatten layer is used to re-shape the inputs from shape [None, 20, 14] to [None, 280] where the first index is the batch size which is flexible and depends on how the data set is prepared.

```
model = keras.Sequential()
model.add(layers.InputLayer(
    input_shape=(n_steps, len(train.columns)-1)))
model.add(layers.Flatten())
model.add(layers.Dense(256))
model.add(layers.Dropout(rate=0.25))
model.add(layers.Dense(64))
model.add(layers.Dropout(rate=0.25))
model.add(layers.Dense(1))
model.summary()
```

It turns out this model performs significantly better than any of the three previous LSTM models. However, the model summary indicates a very large number of model parameters, almost 50 times the number of the basic LSTM model, making this model much more flexible and powerful than the earlier LSTM models:

```

Model: "sequential"

+-----+
| Layer (type)        | Output Shape | Param # |
+=====+=====+=====+
| flatten (Flatten)   | (None, 280)  | 0         |
| dense (Dense)       | (None, 256)  | 71936    |
| dropout (Dropout)   | (None, 256)  | 0         |
| dense_1 (Dense)     | (None, 64)   | 16448    |
| dropout_1 (Dropout) | (None, 64)   | 0         |
| dense_2 (Dense)     | (None, 1)   | 65        |
+=====+=====+=====+
Total params: 88449 (345.50 KB)
Trainable params: 88449 (345.50 KB)
Non-trainable params: 0 (0.00 Byte)

```

15.8 Next Activity Prediction in Business Processes

As another example of a sequence data problem, consider predicting the next activity in a running instance of a business process. In particular, train a model to predict the next activity from a prefix sequence of 5 activities that have already occurred. This is a classification problem, where the class of the next activity is to be predicted.

Complete implementations for this and the other examples in this chapter are available in the following GitHub repo: <https://github.com/jevermann/busi4720-ml>

The project can be cloned from this URL: <https://github.com/jevermann/busi4720-ml.git>

Download the example event log here^a: https://evermann.ca/busi4720/BPI_Challenge_2012.xes.gz

^aThe example log file is taken from the [4TU repository](#) where it has been published under the [4TU general terms of use](#).

First, import the required libraries and use PM4Py to read the event log:

```

import numpy
from tensorflow import keras
from keras import layers
import pandas as pd
import pm4py

# Length of sequences to predict from
prefix_len= 5

# Read the log
log = pm4py.read_xes('BPI_Challenge_2012.xes.gz')

```

Preprocess the data by fixing data types after the import and filtering the log for activity

completion events:

```
log['time:timestamp'] = pd.to_datetime(log['time:timestamp'], utc=True)
log['case:REG_DATE'] = pd.to_datetime(log['case:REG_DATE'], utc=True)
log['case:AMOUNT_REQ'] = pd.to_numeric(log['case:AMOUNT_REQ'])
log['org:resource'] = log['org:resource'].astype(str)

# Retain only activity completion events
log = log[log['lifecycle:transition'] == 'COMPLETE']
```

Filter and sort the log for cases that contain 6 or more activities (5 to predict from, plus one target):

```
# Find the case start time as time of the first event in case
log = log.merge(log.groupby('case:concept:name', as_index=False) \
    ['time:timestamp'].min() \
    .rename(columns={'time:timestamp':'case:start'}), how='left')

# Find the number of events for each case
log = log.merge(log.groupby('case:concept:name', as_index=False) \
    ['time:timestamp'].count() \
    .rename(columns={'time:timestamp':'num_events'}), how='left')

# Filter log for minimum 6 events (5 input, 1 target)
log = log[log['num_events'] > prefix_len]
# Sort log by case start, then by event time
log.sort_values(['case:start', 'time:timestamp'], inplace=True)
```

Next, specify the feature column:

```
# This is the feature to predict (from)
f_name = 'concept:name'
```

Next, identify the "vocabulary", that is, the set of unique activity names for inputs and targets. Rather than converting this to integers using a Keras preprocessing layer, the following code block builds dictionary objects for converting activity names to and from integers. Note that a special end-of-case marker is added to the set of activity names.

```
# Vocabulary
vocab = list(log[f_name].unique()) + ['EOC']
v_size = len(vocab)

# Dictionaries to convert to/from integers
f2int = dict([(s, vocab.index(s)) for s in vocab])
int2f = dict([(k, v) for (k, v) in f2int.items()])
```

The following code block creates a list of activity for each case. This is done by grouping the data frame by case ID ("case:concept:name") and then using the `apply(list)` function to the feature columns of the grouped data frame and naming the resulting column `features`:

```
# Make sequences of feature names for each case
features = log.groupby(['case:concept:name'])['f_name'] \
    .apply(list).reset_index(name='features')
```

The end-of-case marker is added to each list just created and the lists are then converted to lists of integers using the dictionary built earlier:

```
sequences=[l+[ 'EOC'] for l in list(features['features'])]
sequences=[[f2int[i] for i in seq] for seq in sequences]
```

Split sequences into prefix and target using a sliding window over each sequence:

```
data = pd.DataFrame([(seq[i:i+prefix_len], \
    seq[i+prefix_len]) for seq in sequences \
    for i in range(len(seq)-prefix_len)])
```

Then, split the sequence elements into their own data frame columns, that is Pandas Series:

```
# Split the lists into dataframe columns
data = data.assign(**data[0].apply(pd.Series).add_prefix('index_'))
```

Treating each prefix as independent, split the data randomly into a training and validation set.

```
# Divide into train and test set
train = data.sample(frac=0.8)
valid = data.drop(train.index)
```

Then, separate the features from the targets:

```
# Separate X and Y
train_x = train.iloc[:,2:]
train_y = train.iloc[:,1]
valid_x = valid.iloc[:,2:]
valid_y = valid.iloc[:,1]
```

The neural network model is a single-layer stateless LSTM with a Seq2Vec architecture. The input layer specifies the input shape as 5 sequence steps with an undetermined feature size. The features, that is, the activity numbers, are passed to a word embedding layer with 16 output dimensions. That is, each activity number is transformed into a set of 16 values. Note that this occurs for each of the 5 sequence steps fed into the network. Each embedding output is then provided as the input feature vector to the LSTM layer. The LSTM output is passed to a fully-connected output layer with as many units as there are different activity names. The activation function is a softmax function so that this layer returns class membership probabilities.

```
model = keras.Sequential()
model.add(layers.InputLayer(input_shape=(5,)))
model.add(layers.Embedding(input_dim=v_size, output_dim=16))
model.add(layers.LSTM(units=32,
    return_sequences=False, return_state=False, stateful=False))
model.add(layers.Dense(v_size, activation='softmax'))
```

Finally, the model is compiled and trained for 25 epochs:

```
# Compile with loss and optimizer
model.compile(loss='sparse_categorical_crossentropy',
    optimizer='Adagrad',
    metrics=['sparse_categorical_accuracy'])

# Train the model and validate on
model.fit(train_x, train_y,
    validation_data=(valid_x, valid_y),
    epochs=25, shuffle=True)
```

The model summary is shown below, as well as the training progress information for the first and last epoch. The prediction accuracy improves significantly from the first epoch, for both the training and the validation dataset. Additionally, the training and validation losses are similar, suggesting the model does not overfit.

Layer (type)	Output Shape	Param #
embedding (Embedding)	(None, 5, 16)	384
lstm (LSTM)	(None, 32)	6272
dense (Dense)	(None, 24)	792
<hr/>		
Total params: 7448 (29.09 KB)		
Trainable params: 7448 (29.09 KB)		
Non-trainable params: 0 (0.00 Byte)		
Epoch 1/25		
2875/2875 [=====] - 7s 2ms/step -		
loss: 3.0962 - sparse_categorical_accuracy: 0.1984 -		
val_loss: 3.0310 - val_sparse_categorical_accuracy: 0.2013		
Epoch 25/25		
2875/2875 [=====] - 6s 2ms/step -		
loss: 1.4728 - sparse_categorical_accuracy: 0.5886 -		
val_loss: 1.4553 - val_sparse_categorical_accuracy: 0.5935		

To predict from the trained model, use the Keras `predict()` function of the fitted model. The following Python code block takes as input a sequence from the training data set, and predicts the class membership probabilities, that is the probabilities over the following process activity. The predicted activity is determined by sampling from the class membership. Alternatively, the most likely class could be chosen. Both options can be combined in that a random choice is made between sampling from the probabilities and choosing the most likely class.

```

input = train_x.iloc[2:3,:].copy()
print(input)

# Take the probabilities of the first entry of the return batch
probs = model.predict(input)[0]

# Either deterministically choose the most probable class
# pred = probs.argmax()
# Better is to randomly sample from the probabilities
pred = numpy.random.choice(a=range(v_size), p=probs)

# Print the result
print(int2f[pred])

```

The chosen activity is added to the end of the input sequence and the prediction is repeated until the end of the case is predicted.

```
# And keep doing this until end-of-case is reached
while int2f[pred] != 'EOC':
    for i in range(4):
        input.iat[0,i] = input.iat[0, i+1]
    input.iat[0,4] = pred
    print(input)
    probs = model.predict(input) [0]
    pred = numpy.random.choice(a=range(v_size), p=probs)
    print(int2f[pred])
```

An example output is shown below. The exact result depends on the specific parameter values of the trained model and the sampling of the class probabilities.

```
      index_0  index_1  index_2  index_3  index_4
4380       9       10       9      15      13
1/1 [=====] - 0s 11ms/step
W_Nabellen incomplete dossiers
      index_0  index_1  index_2  index_3  index_4
4380      10       9      15      13      21
1/1 [=====] - 0s 13ms/step
O_ACCEPTED
      index_0  index_1  index_2  index_3  index_4
4380       9      15      13      21      13
1/1 [=====] - 0s 12ms/step
W_Nabellen offertes
      index_0  index_1  index_2  index_3  index_4
4380      15      13      21      13       9
1/1 [=====] - 0s 13ms/step
W_Valideren aanvraag
      index_0  index_1  index_2  index_3  index_4
4380      13      21      13       9      15
1/1 [=====] - 0s 16ms/step
EOC
```

Hands-On Exercise

Download the [complete Python file](#) and the [example event log](#).

Adapt the network architecture to identify the impact on training and validation performance of the following:

1. Dropouts in the LSTM layer (use the `dropout` option when defining the LSTM layer)
2. GRU instead of LSTM layers (use `layers.GRU()`)
3. Embedding size (originally 16, note: vocabulary size is 23+1)
4. Further training epochs (originally 25)

Comment on your findings and identify the best model.

15.9 Review Questions

Introduction

1. What are Recurrent Neural Networks (RNNs) and what type of data are they particularly suited for?
2. Explain how RNNs maintain a "memory" of previous inputs. What is the role of the internal state in this process?
3. Describe at least two applications of RNNs in natural language processing and explain why RNNs are well-suited for these tasks.
4. Discuss the use of RNNs in business processes. How can they predict outcomes based on previous sequences of actions?
5. Explain the concept of "vanishing gradients" and how it impacts the training of traditional RNNs.
6. Compare and contrast the capabilities of RNNs with those of other types of neural networks, such as convolutional neural networks, in handling sequential data.
7. Analyze how RNNs can be utilized for forecasting in financial markets. What makes RNNs suitable for this type of prediction?

Sequence Models

8. What does it mean to say that RNNs are "recurrent"? How does this characteristic affect their architecture and functionality?
9. Define the following terms and explain their significance in the context of RNNs: Seq2Vec, Vec2Seq, and Seq2Seq.
10. Provide a detailed use case for each of the model types mentioned above (Seq2Vec, Vec2Seq, Seq2Seq), explaining how each model processes data and the type of outputs they generate.
11. What challenges might arise when using the Seq2Seq model for machine translation? Consider both the input and output sequences in your response.
12. How does the architecture of a Seq2Vec model assist in tasks like sentiment analysis from text sequences? Describe the process from input to output.
13. In the context of an RNN, what is the significance of the initial state, and how does it affect the outcome of the sequence processing?
14. Discuss the advantages and potential drawbacks of using a Vec2Seq model for music generation from a fixed-length input vector.
15. What are the challenges associated with training Seq2Seq models, especially when dealing with very long input sequences?

LSTM Cells

16. What are Long Short-Term Memory (LSTM) cells, and why were they developed for use in recurrent neural networks?
17. Provide an example scenario where the unique features of LSTM would be particularly advantageous over standard RNNs.
18. How do LSTMs address the vanishing gradient problem encountered in traditional RNNs?

19. Describe the function of the forget gate in an LSTM cell. How does it affect the cell memory?
20. What impact does the forget gate have on the long-term memory capabilities of the LSTM?
21. Explain the roles of the input and output gates within an LSTM cell. How do they interact with each other to determine the cell's output?
22. What is the significance of the cell memory c_t in LSTMs? How is it different from the hidden state h_t ?
23. Detail the process of updating the cell memory in an LSTM. What equations are involved, and how do they function together to update the memory?
24. Discuss how the output of an LSTM cell is calculated and what role the output gate plays in this process.

GRU Cells

25. What are Gated Recurrent Units (GRU) and how do they simplify the architecture of traditional LSTMs?
26. Describe the two gates used in GRUs and their functions. How do they compare to the three gates in LSTMs?
27. Explain how the update and reset gates in a GRU work together to determine the hidden state of the cell.
28. How do the reset and update gates in a GRU influence the final output of the cell?
29. How does the GRU model decide how much of the past information to forget and how much of the new information to add?
30. Explain the concept of a candidate state in GRUs and how it is integrated into the overall state update.
31. Discuss the potential advantages and limitations of GRUs compared to LSTMs in terms of training efficiency and performance.

Statefulness in RNNs

32. Explain the difference between stateful and stateless RNNs. How does each approach handle the internal state during training?
33. What are the advantages of using a stateful RNN? In what scenarios might it be preferable to a stateless RNN?
34. Give an example of a real-world application where stateful RNNs could provide significant benefits over stateless configurations.
35. How does batch processing differ in stateful vs. stateless RNNs?

Chapter 16

Intepretable Machine Learning

Sources and Further Reading

The material in this chapter is based on the following sources.

Molnar, Christoph: *Interpretable Machine Learning* (2023)

<https://christophm.github.io/interpretable-ml-book/>

(CC BY-NC-SA License)

Christoph Molnar's book is a very readable and fairly comprehensive introduction to interpretable machine learning or explainable artificial intelligence (XAI). It provides only a few mathematical formulas for illustration and explanation and otherwise relies on intuitive and easy-to-understand descriptions and explanations. It points to and uses software for R and Python, but is not focused on code.

SciKit-Learn: <https://scikit-learn.org/>

SciKit-Learn is a comprehensive machine learning framework for Python that also provides some interpretable ML functions. It provides many commonly used as well as advanced functions for classification, regression, and clustering, as well as methods for data preprocessing and model validation and selection.

PyALE: <https://pypi.org/project/PyALE/>

PyALE is a Python package to compute Accumulated Local Effects. It is based on an R package with similar functionality and provides ALE plots for numerical and

categorical features.

LIME: <https://github.com/marcotcr/lime>

LIME is a Python package to compute Local Interpretable Model Explanations (a local model-agnostic method). The package was developed by the authors of the original paper that developed and introduced this method. LIME can be used to explain tabular predictions as well as text- and image-based predictions.

SHAP: <https://shap.readthedocs.io/en/latest/>

SHAP is a Python package to compute Shapley Additive Explanations (a local model-agnostic interpretation method). The package was created by the authors of the paper that initially developed the method. SHAP can provide explanations for tabular, text and image predictions.

16.1 Introduction

Many machine learning models, in particular non-linear, deep neural network models, are too complex for humans to comprehend or grasp. Such models can have millions or even billions of trainable parameters in complex architectures that may include convolutions, LSTM cells, and other complex elements. These kinds of models are known as "black box" models, because their inner workings are essentially inscrutable.

Interpretable machine learning refers to the techniques and models where the processes and results they produce can be understood by human beings. The aim of interpretable machine learning is to ensure human understanding of how the machine learning models work and how they arrive at their results, that is, their decisions or predictions. In other words, it is about *why* a decision or prediction is made, in understandable terms.

Motivation

This interpretability is crucial for various aspects from debugging models to compliance with legal standards. Interpretable machine learning is important for a number of reasons:

- *Curiosity*: Interpretable models satisfy human curiosity about how decisions are made, especially in systems affecting our daily lives. Understanding the decision process can foster deeper insights and innovations.
- *Human Learning*: By interpreting machine learning models, humans can learn new patterns and knowledge about the data and the phenomena they model, potentially leading to academic advancements and practical applications.
- *Human Sensemaking of Events and Phenomena*: Interpretability helps users make sense of automated decisions and the events or actions that are conse-

quences of such decisions, ensuring that outcomes are understandable in human terms.

- *Knowledge Extraction for Scientific Progress:* Interpretable models allow scientists and researchers to extract and verify new knowledge from complex datasets, advancing scientific understanding.
- *Safety and Compliance Assessment:* Ensuring that machine learning systems operate safely and within regulatory frameworks is easier when these systems are interpretable. Stakeholders can verify compliance through understandable model outputs.
- *Reliability and Robustness Evaluation:* Interpretable models facilitate the evaluation of system reliability and robustness by making it possible to assess how decisions are made under various conditions.
- *Identify Knowledge Limits:* Interpretable models help identify the limits of the knowledge embedded in the models, including areas where the model may lack information or where it is unreliable.
- *Auditability:* Being able to audit machine learning processes transparently helps in maintaining accountability, particularly in sectors like finance and healthcare.
- *Bias Detection & Ensuring Fairness:* Interpretability is key in detecting biases in model predictions and ensuring that machine learning applications are fair and do not perpetuate or exacerbate existing inequalities.
- *Trust and Acceptance:* Users are more likely to trust and accept machine learning solutions that they can understand and feel confident that they are fair and effective.
- *Debugging & Failure Analysis:* When a model's decisions can be traced and understood, identifying errors and the reasons for failures becomes feasible, allowing for more effective debugging and correction.
- *Legal Obligations ("Right to Explanation"):* In many jurisdictions, regulations such as GDPR mandate that decisions made by automated systems be explainable to affected individuals, fulfilling legal obligations.

Intrinsic and Post-Hoc Interpretability

Interpretability in machine learning can be broadly classified into two categories: intrinsic interpretability and post-hoc interpretability. These two types represent different approaches and philosophies in making machine learning models understandable to humans.

Intrinsic interpretability refers to the use of machine learning models that are naturally understandable due to their simple structure and the simplicity of their decision-making process. These models are known as "white box" models and are designed to be interpretable from the ground up, with the following important characteristics:

- *Simplicity*: Models such as linear regression, decision trees, or logistic regression are considered intrinsically interpretable because their decisions can be easily traced to their input features and understood without requiring additional tools or techniques.
- *Transparency*: The model's workings are transparent, meaning that each step of the decision process is visible and comprehensible to users.
- *Limited Complexity*: These models typically involve fewer parameters and simpler relationships, which aids in direct human comprehension.

Post-hoc interpretability involves techniques and methods applied after model training to explain or clarify how a model makes decisions. This is often necessary for complex models like neural networks or ensemble methods that combine multiple classifiers, where intrinsic interpretability is not feasible. Methods such as LIME (Local Interpretable Model-agnostic Explanations), SHAP (SHapley Additive exPlanations), or feature importance metrics are commonly used to provide insights into model decisions. These methods do not depend on the model type; they can be applied to any model type. Post-hoc methods are designed to be used externally to the original black box model, making them versatile in application and providing flexibility in model choice and deployment.

The choice between intrinsic and post-hoc interpretability depends on the balance between the need for prediction accuracy and the requirement for transparency or understandability. While intrinsic interpretability is preferable for clarity and ease of understanding, the simplicity and limited complexity often (but not always) means that these models have lower predictive performance. Post-hoc interpretability provides critical insights into complex models, ensuring that even the most sophisticated algorithms can be scrutinized and understood. However, the understanding is often limited to the output only, rather than the method by which a model generates output. That is, the interpreted model remains a somewhat opaque black box model.

Local and Global Interpretation

Interpretation methods are often categorized into local and global methods. Each type provides different insights into the functioning of machine learning models, tailored to specific needs and scenarios.

Local interpretation methods focus on explaining individual predictions made by a machine learning model. These methods aim to clarify why a model made a specific classification decision or regression prediction for a particular instance. These methods provide explanations for individual data points, helping to understand the model's behavior at a granular level. Examples include LIME (Local Interpretable Model-agnostic Explanations) and counterfactual explanations, which explain how changes in input features could alter the prediction. Local interpretations are particularly useful in applications like healthcare or finance, where understanding specific decisions is crucial for trust and verification. Typically, local methods focus on the impact of specific feature values.

Global interpretation methods seek to explain the overall behavior or logic of a model across all instances. They provide a holistic view of how the model operates in general. These methods explain the model's general decision-making process rather than focusing on individual predictions. Examples include feature importance and partial dependence plots, which show the effect of each feature on the model's predictions across the entire dataset. Global interpretations are beneficial when stakeholders need to validate the model's overall logic and ensure it aligns with domain knowledge and objectives. In contrast to local methods, global methods focus on the impact of features, not the impact of specific feature values.

The choice between local and global interpretation methods depends on the specific needs for transparency in a given application. Local methods are best suited for cases requiring detailed explanations of individual decisions, while global methods are ideal for understanding and validating the overall behavior of a model.

16.2 Intrinsically Interpretable Models

Intrinsically interpretable machine learning models often leverage characteristics such as linearity, monotonicity, and interactions to ensure that their workings are transparent and understandable.

Linearity in machine learning models means that the output is a linear combination of the input features. Models like linear regression are classic examples where each predictor has a constant effect on the outcome. Linear models are straightforward, making them easy to understand and explain. The impact of each feature on the prediction is clear and quantifiable. The effects of changes in input features are predictable, aiding in scenario analysis and planning.

Monotonicity in a model ensures that the relationship between any given input and the output is either always non-decreasing or always non-increasing. Linear regression models and decision trees that split based on a single feature at a time can exhibit this property. Monotonic models are consistent in their behavior, enhancing trust as increases (or decreases) in input variables lead to predictable changes in outputs. These models are easier to validate against domain knowledge, where certain inputs are expected to have a direct and consistent influence on the output.

Interactions in machine learning refer to the scenario where the effect of one feature on the response variable depends on the value of another feature. By modeling the interdependencies between features, interaction-enabled models can often achieve higher predictive accuracy while still maintaining a degree of interpretability. Many phenomena involve interdependent factors; accurately modeling these interactions can make the model outputs more applicable and relevant in real-world scenarios.

Linearity, monotonicity, and interactions each play crucial roles in ensuring that intrinsically interpretable machine learning models are both effective and transparent. While linearity and monotonicity contribute to simplicity and predictability, interactions allow for a nuanced understanding of complex dynamics within the data. Table 16.1 shows a

Algorithm	Linear	Monotone	Interaction
Linear regression	Yes	Yes	No
Logistic regression	No	Yes	No
Decision trees	No	Some	Yes
RuleFit	Yes	No	Yes
Naive Bayes	No	Yes	No
k-NN	No	No	No

Source: <https://christophm.github.io/interpretable-ml-book/simple.html>

Table 16.1: Intrinsically Interpretable Models

list of intrinsically interpretable models. This section examines the emphasized entries in the table, linear regression and decision trees, while others, such as logistic regression, naive Bayes, and k-NN, have been presented and discussed in previous sections.

16.2.1 Linear Regression

Linear regression is one of the simplest prediction methods available and is intrinsically interpretable. Consider the following example linear regression model for predicting bicycle rental count (variable `cnt`) from the season (`season`) and the temperature (`cnt`).

```
# Load the bike rental data set
d <- read.csv('https://evermann.ca/busi4720/bike.csv')

# Perform the regression and summarize results
summary(lm(cnt ~ season + temp, data=d))
```

The abbreviated output below is easy to interpret and demonstrates linearity and monotonicity. For example, the coefficient for temperature of 132.79 means that, all other variables remaining the same ("ceteris paribus"), a change of one degree of temperature will change the predicted bicycle rental count by 132.79, no matter what the original temperature is (linearity) and always in the same direction as the temperature change (monotonicity).

The coefficients for the categorical season variable are also easy to interpret. They represent the predicted change in bicycle rental count from the reference category (in this case, FALL), assuming all other variables remain the same ("ceteris paribus"). For example, the prediction for the winter is 1342.87 lower than for the fall.

The intercept is easy to interpret as that prediction when all other variables are 0, or the reference category for categorical variables. In this example, for a temperature of 0 degrees and in the fall season, the model predicts a rental count of 3151.02.

```
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 3151.02    169.35 18.606 < 2e-16 ***
seasonSPRING -494.15    163.28 -3.026 0.00256 **
seasonSUMMER -852.68    209.82 -4.064 5.35e-05 ***
seasonWINTER -1342.87   164.59 -8.159 1.49e-15 ***
temp          132.79     11.02 12.046 < 2e-16 ***
---
Residual standard error: 1433 on 726 degrees of freedom
Multiple R-squared: 0.4558, Adjusted R-squared: 0.4528
```

The R^2 is an easily understood metric to assess the quality of the model. It explains the proportion of variance in the target variable that is explained by the predictors. A good model has an R^2 close to 1, while an R^2 close to 0 indicates a poor model.

Finally, the *relative feature importance* is given by the $t = \frac{\hat{\beta}}{SE(\hat{\beta})}$ value. Because it divides the coefficient estimate by its standard error, the t-value is scale invariant and can be used to compare the importance of features on different scales. In the above example, it is clear that temperature is the most important features.

In summary, it is easy to understand which features are most or least important in making a prediction and how features interact or do not interact (consider what "ceteris paribus" implies for interactions).

Linear regression also has high local interpretability; predictions for individual observations are also easy to understand. For example, to predict the bicycle rental count in summer when the temperature is 20 degrees, one uses the following regression equation, with parameter values from the output above:

$$\hat{c} = 3151.02 - 852.68 + 20 \times 132.79 = 4954.14$$

Finally, the ordinary least squares algorithm that is used to compute the model parameters is simple and well-understood and is proven to produce optimal parameter values. This means that linear regression has high *algorithmic transparency*. In other words, it is clear what the optimization objective is, and how it is achieved.

To further improve the interpretability of linear regression models, one may try to reduce the number of features or input dimensions, using methods such as:

- *Manual feature selection*: The analyst may successively omit less important features, or may include features based on the strength of their correlation with the target variable.
- *Automatic feature selection*: While an exhaustive search of all possible feature combinations is typically not feasible, automated heuristics are able to arrive at a reasonably good subset of features that still provides good prediction capability with a small number of features.

- *Regression with PCA components:* PCA produces linear combinations of variables with maximal variance. Retaining a few of such components instead of the full set of features may make the model more interpretable. However, the components themselves must also be interpreted, based on their loadings or correlations with the original features.
- *Penalized regression:* The LASSO is a penalized regression that automatically removes features (sets their coefficient to 0) as a function of the penalty parameter λ .

In general, the analyst must consider the bias and variance implications of these dimensionality reduction methods: Simpler models will generally have a higher bias and lower variance than more complex models.

16.2.2 Decision Trees

Decision trees are another relatively simple and easy to understand predictive machine learning technique. Decision trees can be used both for regression ("regression trees") and for classification ("classification trees"). Decision trees are a popular choice in interpretable machine learning due to their strengths:

- *Algorithmic Transparency:* The procedure to construct a decision tree and the optimization criteria are easy to understand and verify. The optimization criterion to derive the decision rules is simple and intuitive.
- *Simplicity:* The structure of a decision tree is simple and intuitive, which allows non-experts to understand the model's reasoning easily. This simplicity also facilitates easier communication about the model's decision-making process.
- *Visualization:* Decision trees offer a clear visualization of how decisions are made, with each node representing a decision rule and each path a decision process. This makes them highly interpretable and easy to follow.
- *No Need for Feature Scaling:* Unlike some other machine learning models, decision trees do not require input features to be scaled or normalized. This avoids potential distortions in interpretation due to preprocessing.
- *Handling of Non-linear Relationships:* Decision trees can handle complex, non-linear relationships between features without needing any transformation of data, capturing interactions between the variables naturally.
- *Capability to Handle Both Numerical and Categorical Data:* Trees can process datasets that have a mix of numerical and categorical variables without the need for extensive preprocessing.

However, decision trees have a number of weaknesses that may limit their usefulness or applicability to certain problems.

- *Overfitting:* Decision trees are prone to overfitting, especially for very deep trees. Without aggressive tree "pruning", they can create overly complex trees that do not generalize well.

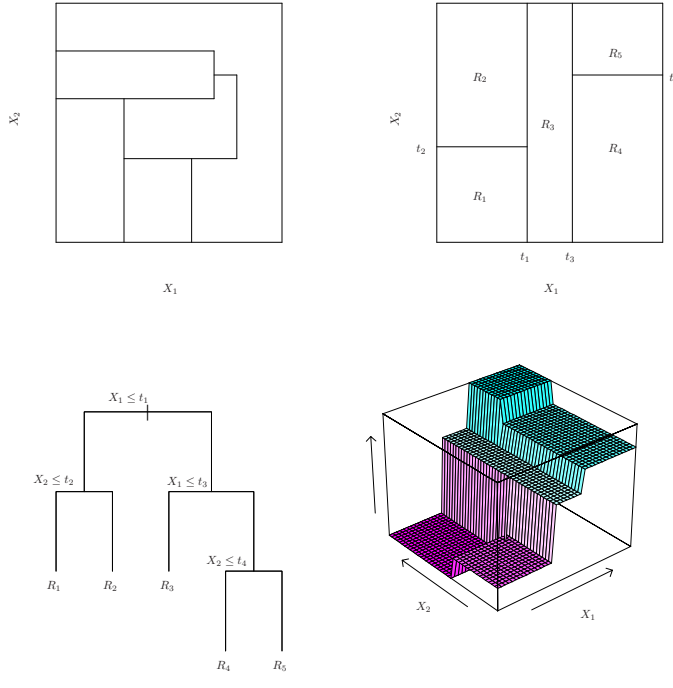


Figure 16.1: Decision tree for two numerical features

- *Instability:* Small changes in the data can result in a completely different tree being generated. In other words, they have high variance.
- *Bias Towards Certain Structures:* Decision trees tend to favor splits on features having a larger number of levels. This can bias the model's decision-making process, especially if the feature's relevance is artificially inflated by its structure.
- *Greedy Nature:* Standard algorithms for decision tree construction, such as CART (Classification and Regression Trees, discussed below), are greedy because they make the best split at a given node without considering future impacts. This might lead to suboptimal trees.
- *Piecewise Constant Predictions:* Decision trees make piecewise constant decisions, that is, they predict the same values or classes within small regions of the feature space. This means that the predictions within these spaces have zero variance, which may be problematic for subsequent analytics.

Figure 16.1 shows the intuition behind binary decision trees. The top left panel in the figure shows a feature space spanned by two numerical features, x_1 and x_2 . The space is subdivided into regions where predictions on the target variable are similar. Unfortunately, an optimal sub-dividing of the feature space is difficult to achieve when

the regions are irregularly shaped. Instead, binary decision trees are created by making a binary split of the feature space along one feature/dimension, yielding two smaller regions. Each region may then be split again, recursively, until a stopping criterion is reached.

Consider the top right panel in Figure 16.1 as an example. Here, the feature space is first split on feature x_1 at value t_1 creating two rectangular regions, one to the left of t_1 where $x_1 \leq t_1$ and the other to the right where $x_1 > t_1$. The left region is then split on feature x_2 at value t_2 creating the regions labelled R_1 where $x_1 \leq t_1 \wedge x_2 \leq t_2$ and R_2 where $x_1 \leq t_1 \wedge x_2 > t_2$. These two regions are not split any further, and are called the "leafs" or "leaf nodes" of the tree. The region to the right of t_1 , that is the region where $x_1 > t_1$, is split again on feature x_1 at value t_3 , yielding the region R_3 where $x_1 > t_1 \wedge x_1 \leq t_3$ and the region where $x_1 > t_1 \wedge x_1 > t_3$. The latter region is split on feature x_2 at value t_4 , yielding regions R_4 and R_5 .

The bottom left panel in Figure 16.1 shows how these split or divisions of the feature space form a (upside down) decision tree with root, decision nodes, and the resulting leaf nodes. The bottom right panel in Figure 16.1 shows what a prediction of a target variable (here, a numerical target), might look like. For a regression tree, the predicted value for each region is simply the average of the observed target values in each region, leading to the flat plateaus shown in the bottom right panel, that is, piece-wise constant predictions for small regions of the feature space. For a classification tree, the predicted class for a leaf node is simply the majority class of observations in each region.

For binary decision trees, the choice on which feature to split on next and which specific feature value to split on is decided as follows:

1. For every feature j and potential split feature value s define regions

$$R_1(j, s) = \{X | X_j < s\} \quad \text{and} \quad R_2(j, s) = \{X | X_j \geq s\}$$

This is simple for binary feature but requires dichotomization or discretization (for example, using equal interval bins or equal frequency bins) for numerical features and can potentially result in a large number of possible split points s .

2. Use one of the following criteria to choose j and s :

- *Minimize the total variance of the two regions (regression trees):*

$$\sum_{i:x_i \in R_1(j,s)} (y_i - \bar{y}_{R_1})^2 + \sum_{i:x_i \in R_2(j,s)} (y_i - \bar{y}_{R_2})^2$$

The decision tree algorithm will calculate the total variance of both groups resulting from each potential split and then choose the split that will yield the lowest variance.

- *Maximize information gain (classification trees):* Recall that the entropy is defined as

$$H(Y) = - \sum_{y \in Y} p(y) \log p(y)$$

where $p(y)$ represents the proportion of each class y in a child node. The information gain obtained by splitting on a feature j at feature value s and forming regions R_1 and R_2 is then defined as:

$$IG(Y, a) = H(Y) - p(R_1)H(Y|R_1) - p(R_2)H(Y|R_2)$$

where $p(R_1)$ is the probability of an observation being in region R_1 , that is, the proportion of observations in region R_1 , and $H(Y|R_1)$ is the conditional entropy, that is, the entropy of those observations in region R_1 (analogous for R_2).

Intuitively, the entropy H expresses the uncertainty in the distribution of Y . Knowing or assuming a value (or range of values) for some feature j should decrease the uncertainty, that is, lead to a gain in information. The decision tree algorithm will calculate the information gain resulting from each split and will choose the split with the highest gain.

- *Minimize Gini index (classification trees):* The Gini impurity of a dataset is defined as:

$$Gini = 1 - \sum_{i=1}^k p(y)^2$$

where $p(y)$ is the probability of an observation being classified to a particular class y . The Gini impurity index represents the probability of an observation being misclassified if it were randomly assigned a label according to the distribution of classes in the data at that node. A Gini impurity of 0 indicates that all cases in the node fall into a single category, i.e., the node is completely pure. The decision tree algorithm will calculate the Gini impurity for each subgroup that would result from a split and will choose the split that results in the lowest weighted sum of the group impurities.

The above process is recursively iterated for each region, resulting in a tree that is growing new leaf nodes and gaining depth. Without any stopping, the tree will grow, that is, nodes will be split, until every leaf node contains only a single observations. It is clear that such a fully-grown tree is overfitted to the training data set, and not easily interpretable.

In practice, to prevent overfitting and ensure the tree remains interpretable, the splitting process is stopped using one many options for *stopping criteria*:

- *Maximum tree depth:* The splitting of the feature space stops when a specified tree depth is reached. All branches of the tree have the same depth/length.
- *Maximum leaf node count:* Splitting of the feature space stops when a certain number of leaf nodes have been created. Not all branches of the tree need to have the same length/depth. This limits the number of decisions in the decision tree and the number of unique predicted values for regression trees.

- *Minimum Leaf node sample size*: Leaf nodes must have a minimum number of observations. This makes the predictions for each feature space region more representative as they are based on a minimum sample size.
- *Maximum leaf node variance*: Splitting continues until the variance of observations in a regression tree regions is below a certain threshold.
- *Minimum leaf node purity*: Regions are split while they contain a mixture of target observations; splitting stops only when a node contains a certain proportion of a majority class for classification trees or when the entropy is below a certain threshold. This criterion prevents unnecessary splitting which may result in two sub-regions yielding the same predicted class.
- *Minimum change in node impurity or information gain*: Regions are split while the improvement in purity or reduction of entropy are above a certain threshold. This criterion prevents unnecessary splits that do not significantly improve the predictive power of the model.

To illustrate decision trees, consider the following Python example, using the `DecisionTreeRegressor` class from the SciKit-Learn package. The example uses the same data set as the linear regression example above and constructs a regression tree to predict the count of bicycle rentals from the temperature and humidity, two numerical variables.

The example fits an unpruned tree, that is, there is no stopping criterion and the tree will have as many leaf nodes as there are observations. Consequently, there will be no prediction error for the training data and the tree is very much overfitted.

```
import matplotlib.pyplot as plt
import pandas as pd
# Prepare data:
d=pd.read_csv('https://evermann.ca/buci4720/bike.csv')
x=d[['temp', 'hum']]
y=d['cnt']
#Fit unpruned tree:
from sklearn.tree import DecisionTreeRegressor
regr = DecisionTreeRegressor()
regr.fit(x, y)
# Print the MSE:
from sklearn.metrics import mean_squared_error
mean_squared_error(regr.predict(x), y)
```

Early stopping can prevent overfitting and maintain interpretability. The following three examples show how to use the tree depth criterion, the leaf node sample size criterion and the leaf node count criterion for stopping the tree construction. Each of those trees will have a non-zero training error.

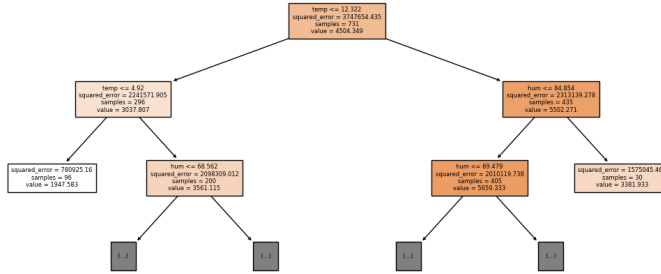


Figure 16.2: Regression tree example

```

regr = DecisionTreeRegressor(max_depth=3)
regr.fit(x, y)
regr = DecisionTreeRegressor(min_samples_leaf=10)
regr.fit(x, y)
regr = DecisionTreeRegressor(max_leaf_nodes=8)
regr.fit(x, y)
  
```

To demonstrate the interpretability of a decision tree, it is useful to print the tree and to visualize or plot the tree.

```

import sklearn
# Print the tree:
print (sklearn.tree.export_text(regr, feature_names=x.columns))
# Plot the tree:
sklearn.tree.plot_tree(regr, max_depth=2, feature_names=x.columns,
                      filled=True, fontsize=6)
plt.show()
  
```

Printing the fitted tree will show the decision nodes and the predicted value of each leaf node, which is simply the mean value of the observations for that leaf node. The output from the Python code above is shown below. Plotting the tree results in the visualization shown in Figure 16.2. It is clear that decision trees have good local interpretability, that is, for a given observation, it is easy to see how the tree model arrives at its predictions. However, global interpretability is lacking. For example, it is unclear whether temperature or humidity is the more important predictor in this example. One could count the number of splits that are based on a particular variable to gain an idea of its importance, but this is a pretty rough approximation.

```

|--- temp <= 12.32
|   |--- temp <= 4.92
|   |   |--- value: [1947.58]
|   |--- temp >  4.92
|   |   |--- hum <= 68.56
|   |   |   |--- value: [3885.62]
|   |   |--- hum >  68.56
|   |   |   |--- value: [2916.96]
|--- temp >  12.32
|   |--- hum <= 84.85
|   |   |--- hum <= 69.48
|   |   |   |--- temp <= 17.40
|   |   |   |   |--- value: [5355.81]
|   |   |   |--- temp >  17.40
|   |   |   |   |--- temp <= 23.51
|   |   |   |   |   |--- value: [6698.34]
|   |   |   |--- temp >  23.51
|   |   |   |   |--- value: [5716.78]
|   |   |--- hum >  69.48
|   |   |   |--- value: [5183.22]
|--- hum >  84.85
|   |--- value: [3381.93]

```

Finally, to show the piecewise constant predictions across small regions of the feature space, consider the scatter plot of predicted values in Figure 16.3, created using the following Python code block. The plot shows the actual target on the horizontal axis and the predicted target on the vertical axis. Predictions for different actual target values are the same. From the graph, it is evident that the tree has eight leaf nodes corresponding to those in the printed output, that is, the tree is able to predict 8 different values.

```

# Plot fitted versus true values:
import plotly.express as px
px.scatter(pd.DataFrame([y, regr.predict(x)], \
    index=['y', 'yhat']).transpose(), x='y', y='yhat').show()

```

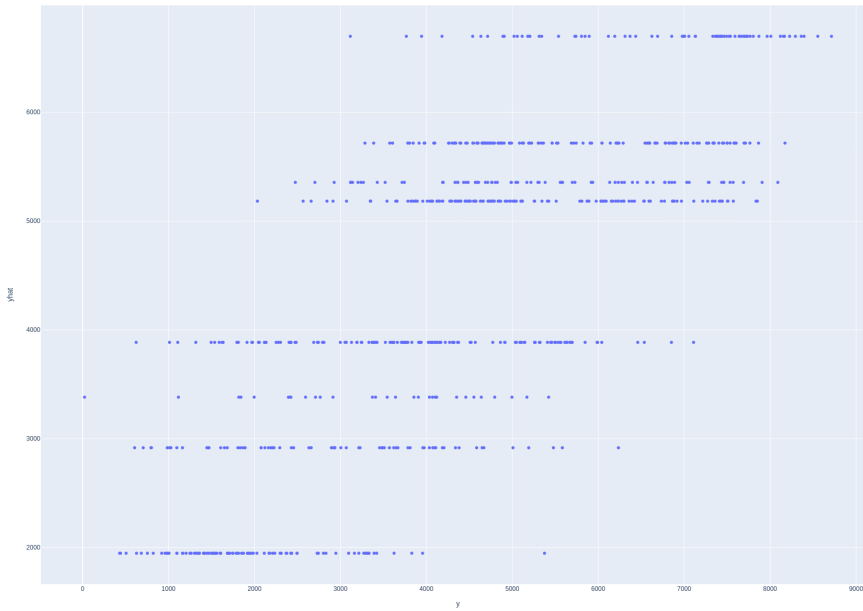


Figure 16.3: Predictions of a regression tree

Hands-On Exercise

1. Fit regression trees to the bike dataset in the above examples. Calculate the MSE, print the decision rules, and plot predicted versus true values as you vary the following stopping rule criteria:

- `max_depth`: choose values 1, 3, 5, 7
- `min_samples_leaf`: choose values 1, 5, 10, 20
- `max_leaf_nodes`: choose values 2, 8, 16, 32

How does the training MSE change? What can you observe from the plots of predicted versus true values?

2. Split the data set into a training and test sample. Repeat exercise (1) but now evaluate the test MSE. What is the optimal stopping criterion?

16.3 Global Model-Agnostic Methods

Global model-agnostic methods focus on the understanding of the overall impact of features in determining the prediction. They are applicable to different models and focus only on the input features and predictions, while ignoring the specific model or algorithm. This section presents a few of the many methods that have been proposed and developed in recent years.

16.3.1 Partial Dependence Plots (PDP)

Partial Dependence Plots (PDPs) provide insights into the relationship between a feature and the predicted outcome, averaged over the distribution of values of the other features in the dataset. By isolating the effect of one or two features, PDPs can show how changes in these features impact the predicted outcome, irrespective of the values of other variables, offering a global view of the model's mechanics.

Formally, a PDP shows the marginal effect of one or a few features X_S on the predicted output, marginalized over all other ("complement") features X_C . Recall that marginalization is the summing, or integration for continuous features, weighted by probabilities:

$$\begin{aligned}\hat{f}_S(X_S) &= \mathbb{E}_{X_C} [\hat{f}(X_S, X_C)] \\ &= \int \hat{f}(X_S, X_C) p(X_C) dX_C\end{aligned}$$

When estimated from sample data, this becomes:

$$\hat{f}_S(X_S) = \frac{1}{n} \sum_{i=1}^n \hat{f}(X_S, X_C^{(i)}) \quad (16.1)$$

where the sum in Equation 16.1 is over all observations and $X_C^{(i)}$ are the complement feature values of the i -th observations. The integrals or sums are computed for all possible values of X_S to generate the PDP curves. Essentially, the PDP shows how the *average* prediction changes when the focal predictor X_S is changed. *Importantly, PDPs assume feature independence.*

Consider the following example in Python. For ease of illustration, the example explains a decision tree model, which is of course intrinsically interpretable already. The example uses the bike rental data set from above and first fits a regression tree.

```
# Read the data set and identify features and target
import pandas as pd
d=pd.read_csv('https://evermann.ca/buci4720/bike.csv')
x=d[['temp', 'hum']]
y=d[['cnt']]
# Fit a regression tree:
from sklearn.tree import DecisionTreeRegressor
regr = DecisionTreeRegressor(max_depth=5).fit(x, y)
```

Once the model is fitted, the `PartialDependenceDisplay` class in the Scikit-Learn package can be used to generate a PDP. The `features` argument indicates which features to use for the PDP. This example Python code below shows individual PDPs for the first and second feature and a joint feature PDP for those two features

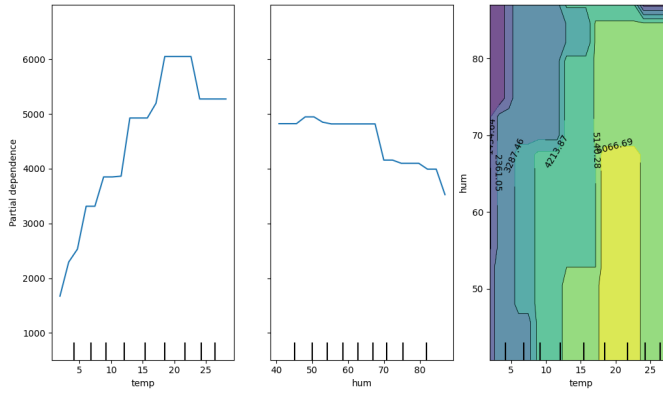


Figure 16.4: Example PDP for a regression tree

as well. The `grid_resolution` argument controls the discretization of continuous features into equal intervals for summing/marginalization. Figure 16.4 shows the resulting PDPs. For example, the left panel shows how the prediction varies when the temperature value changes. For each value of temperature, the sum in Equation 16.1 is computed to calculate the average prediction for that temperature value.

```
import matplotlib.pyplot as plt
from sklearn.inspection import PartialDependenceDisplay
PartialDependenceDisplay.from_estimator(
    estimator = regr,
    X = x, features = [0, 1, (0,1)], grid_resolution = 20)
plt.show()
```

Importantly, the PDP removes the "ceteris paribus" assumption that linear regression models make. Instead of assuming that all other values remain unchanged, the PDP marginalizes ("averages") over the values of other features.

16.3.2 Individual Conditional Expectation (ICE) Curves

Individual Conditional Expectation (ICE) curves extend the idea of PDPs by disaggregating the effects for individual observations. Instead of an average effect over all observations or feature values, an ICE plot displays one curve per instance in the dataset, showing how predictions change for that instance if the feature of interest is varied. This essentially means that no summation in Equation 16.1 takes place; the ICE curves are simply the individual summands of that sum. Thus, averaging all ICE curves yields the PDP curve. This allows identification of outlier cases or heterogeneous data.

ICE plots can be created in Python using the same way as PDPs. The result of the following Python code block is shown in Figure 16.5. While the bike rental data set in this example has 731 observations, the ICE plot in Figure 16.5 shows fewer lines because

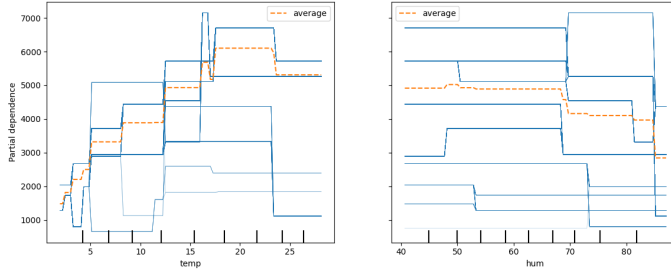


Figure 16.5: Example ICE plot for a regression tree

of the piece-wise constant predictions (the fitted tree has 31 leaf nodes), that is, the predictions for multiple observations are identical and the lines are overlaid on top of each other in the ICE plot. From Figure 16.5 it is clear that not all observations follow the same, average pattern and that there is significant heterogeneity in the data set.

```
PartialDependenceDisplay.from_estimator(regr, x, [0, 1], kind='both')
```

16.3.3 Accumulated Local Effects (ALE) Plot

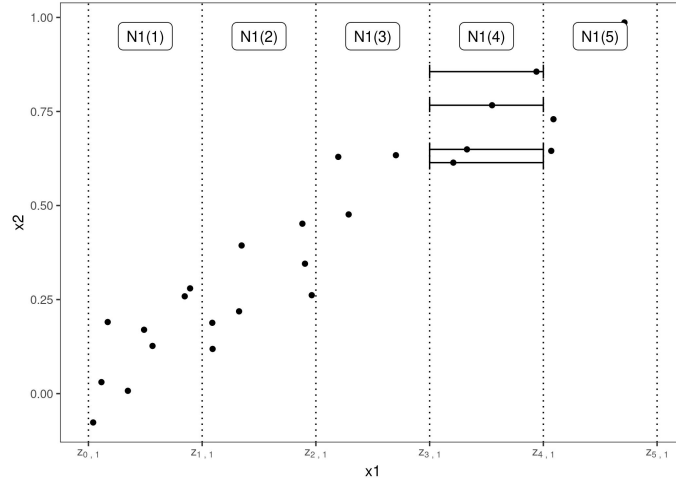
Accumulated Local Effects (ALE) plots address some limitations of PDPs by focusing on local rather than global average effects of features. ALE plots calculate the contributions of features to predictions by accumulating localized average partial effects, helping to avoid misleading interpretations in the presence of correlated features.

Specifically, ALE plots do not construct unrealistic feature combinations as they average over the complement feature values. For example, the temperature and humidity features are mildly correlated in the running example, and not all combinations are realistic. Instead, effects are computed for a grid of intervals, that is, a "local window", instead of the entire domain of a features, as in PDPs.

Formally, the ALE is defined as:

$$\hat{f}_{j,ALE}(X) = \sum_{k=1}^{k_j(x)} \frac{1}{n_j(k)} \sum_{i: x_j^{(i)} \in N_j(k)} \left[\hat{f}(z_{k,j}, x_j^{(i)}) - \hat{f}(z_{k-1,j}, x_j^{(i)}) \right] \quad (16.2)$$

The range of values for feature j is divided into k equal size intervals. $N_j(k)$ is the k -th such local neighbourhood for a feature j with $Z_{k,j}$ being the lower boundary of that interval. The summands in the inner sum of Equation 16.2 are differences in prediction between the lower and upper boundary of a local neighbourhood (specifically, between the lower bound of interval $k-1$ and that of the next interval k). The inner sum is taken



Source: Molnar, Fig. 8.7

Figure 16.6: Illustrative example of ALE computation

over all observations in the neighbourhood. This is termed the "local effect". The outer sum in Equation 16.2 is taken over all local neighbourhoods k , each one weighed by the number of observations $n_j(k)$ in that neighbourhood. In other words, the outer sum *accumulates* the local effects, leading to the ALE.

Consider the illustrative example in Figure 16.6. It shows five neighbourhoods defined for feature x_1 . The four horizontal lines represent the differences in prediction for four observations when the value of x_1 changes from $Z_{3,1}$ to $Z_{4,1}$. The value of x_2 is held constant. The four differences are summed to yield the local effect. Local effects are calculated in the same way for the other four intervals in Figure 16.6 and then summed.

To illustrate ALEs, consider the following regression tree, again trained on the bicycle rental count data with the same features and predictors as before.

```
# Train model:
from sklearn.tree import DecisionTreeRegressor
regr=DecisionTreeRegressor(min_samples_leaf=10).fit(x,y)
```

The PyALE package provides the `ale` class to calculate the accumulated local effects. They can be plotted for a single feature or for two features. Figure 16.7 shows the ALE for the temperature feature on bicycle rental counts. The effect of temperature varies across the range of temperature. The two-feature ALE plot is shown in Figure 16.8 and this also shows a variable impact of the feature combinations on the predicted bicycle rental count.

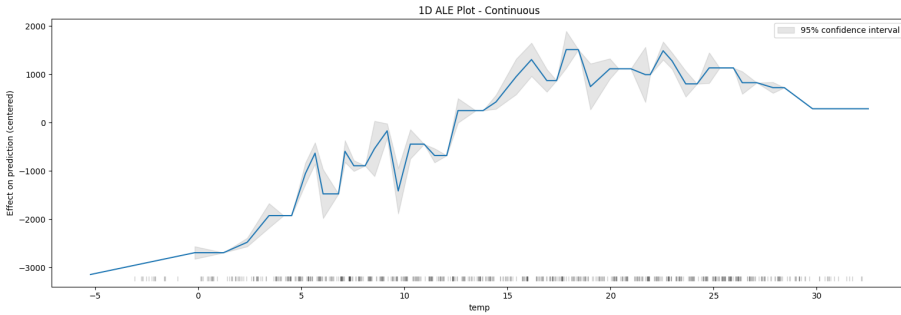


Figure 16.7: Example ALE for one feature

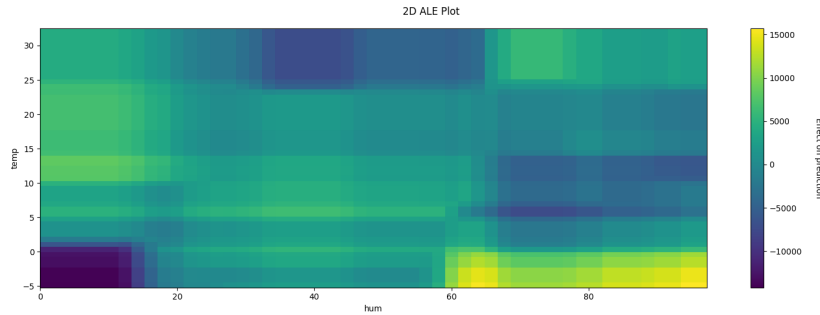


Figure 16.8: Example ALE for two features

```

import matplotlib.pyplot as plt
from PyALE import ale
# Construct the ALE and plot:
ale_effects = ale(X=x, model=regr,
                    feature=['temp'], grid_size=50)
plt.show()
ale_effects = ale(X=x, model=regr,
                    feature=['temp', 'hum'], grid_size=50)
plt.show()

```

Notably, ALE plots also remove the "ceteris paribus" assumption that all other values remain equal. However, instead of averaging over all values of the complement features, as PDPs do, the effects are computed for a local window of complement feature values. This makes ALE suitable for the use with correlated variables.

16.3.4 Permutation Feature Importance

Permutation feature importance assesses the importance of each feature by observing the effect on predictive accuracy when the values of that feature are randomly shuffled.

This disrupts the relationship between the feature and the outcome, and a significant drop in model performance indicates that the feature is important for model predictions. The intuition is simple: A feature that has no effect on the prediction outcome can be randomly reshuffled without consequence for the predictive performance of the model.

Permutation feature importance begins by estimating the prediction error (loss function) for the model L on the original data X . Permutation feature importance should be calculated on test data, not training data.

$$e^{\text{orig}} = L(y, \hat{f}(X))$$

Next, the feature importance is calculated for each feature j , using K multiple random shuffles ("permutations") k of each feature's values:

- For each feature j :
 - For each repetition k in $1 \cdots K$:
 - * Generate permuted data $X_{j,k}^{\text{perm}}$ by permuting ("shuffling") values of feature j
 - * Estimate prediction error of model L for permuted data:

$$e_{j,k}^{\text{perm}} = L(y, \hat{f}(X_{j,k}^{\text{perm}}))$$

- Calculate permutation feature importance:

$$i_j = e^{\text{orig}} - \frac{1}{K} \sum_k^K e_{j,k}^{\text{perm}}$$

The following example again uses the bicycle rental data set. The rental count is used as prediction target, variables other than year and days since 2011 are used as features. A regression tree is fitted with the following Python code block.

```
import pandas as pd
# Prepare data:
d=pd.read_csv('https://evermann.ca/buci4720/bike.csv')
x=pd.get_dummies(d.drop(['yr','days_since_2011'],axis=1))
y=x.pop('cnt')
# Train model:
from sklearn.tree import DecisionTreeRegressor
regr=DecisionTreeRegressor(min_samples_leaf=10).fit(x,y)
```

The SciKit-Learn package provides the `permutation_importance` class to compute feature importance. The `n_repeats` parameter specifies the number of permutations to use for each feature. For simplicity of exposition, the permutation feature

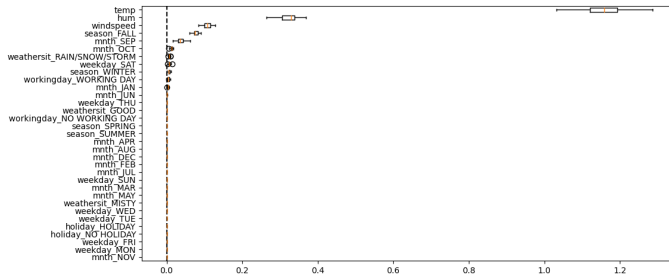


Figure 16.9: Permutation feature importance plot. Uncertainty represents variation of multiple permutations

importance is calculated here using the training data, but practical applications should use independent test data for this. The second line in the code below sorts the permutation feature importances by their mean for later plotting.

```
from sklearn.inspection import permutation_importance
# Calculate permutation feature importance and sort them
r = permutation_importance(regr, x, y, n_repeats=30)
r_idx = r.importances_mean.argsort()
```

The feature importances can be easily visualized using the following Python code. The resulting graph is shown in Figure 16.9. The graph shows that temperature has the greatest importance, followed by humidity and windspeed. Many features have little or no impact on the prediction outcome.

```
import matplotlib.pyplot as plt
# Produce a plot of sorted feature importance:
fig, ax = plt.subplots()
ax.boxplot(r.importances[r_idx].T, vert=False, labels=x.columns[r_idx])
ax.axvline(x=0, color="k", linestyle="--")
plt.show()
```

16.3.5 Global Surrogate Models

Global surrogate models are interpretable models that are trained to approximate the predictions of a complex model. By training a simpler model (like a decision tree or linear regression model) to mimic the behavior of a more complex model, insights can be gained into how the complex model makes decisions, providing an interpretable approximation of its decision-making process. Intuitively, the predictions of the complex model are the prediction targets of the simpler surrogate model, that is, the simple surrogate model should predict the predictions (rather than the original target values).

PDP/ICE	
Intuitive	Limited number of features
Clear interpretation	Assumes feature independence
Easy to implement	
ALE	
Unbiased for correlated features	Local interpretation only
Clear interpretation	ALE may differ from linear coefficients
Faster to compute than PDP	No ICE curves Unstable for large number of intervals
PFI	
Clear interpretation	Linked to model error
Concise, global measure	Requires access to true targets
Does not require retraining	May be biased for correlated features
Takes into account all interactions	
Global Surrogate Models	
Flexible	Conclusions about model, not data
Intuitive	Unclear cut-off for goodness of fit
R-squared measure for fit	

Table 16.2: Strengths and weaknesses of different global model-agnostic methods

Doing this in practice is simple. For example, the following Python code trains a neural network regression model with two hidden layers and ReLU activation function on the bicycle rental data set. The model is then used to predict values for the training observations.

```
# Example 'black box' model
from sklearn.neural_network import MLPRegressor
regr = MLPRegressor((4, 2,), max_iter=10000)
regr.fit(x, y)
preds = regr.predict(x)
```

In the second step, a simpler surrogate model is used to predict the predictions. The following example uses a linear regression model fitted using ordinary least squares. As noted above, the absolute value of the t statistic in the fitted OLS model serves as an indication of feature importance in this surrogate model. The summary results indicate that the surrogate model has an R^2 value of 0.995, that is, it explains the predictions of the complex model (not the original targets!) very well.

```
from statsmodels.api import OLS
OLS(preds, x).fit().summary()
```

Table 16.2 summarizes the strengths and weaknesses of the different global model-agnostic methods for interpretable machine learning discussed in this section.

16.4 Local Model-Agnostic Interpretation Methods

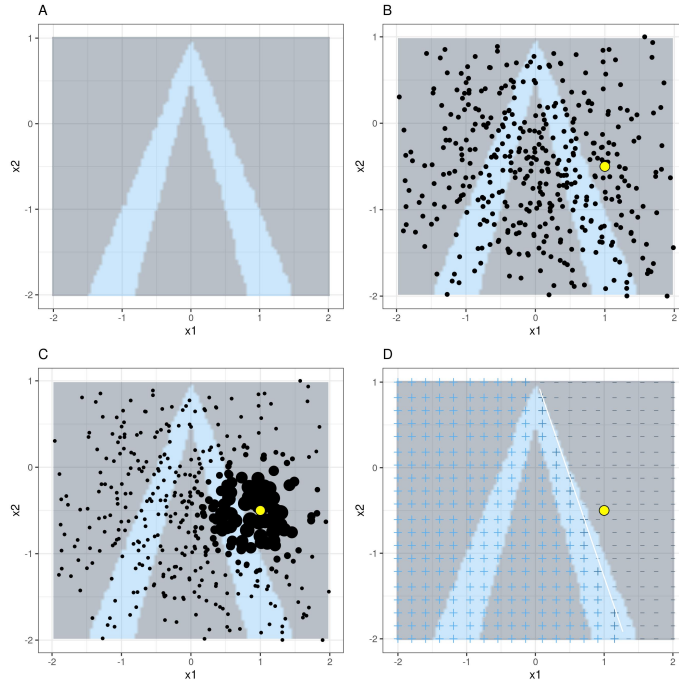
Local model-agnostic interpretable methods aim to provide explanations for individual predictions or cases, regardless of the overall model's complexity or opacity, offering insights into specific decision points rather than a model's global behavior. This section presents two widely-used methods that each address the challenge of interpretability from a different perspective.

16.4.1 Local Interpretable Model-agnostic Explanations (LIME)

The intuition behind LIME is that while it may be difficult or impossible to explain an entire complex model globally, it is possible to approximate and explain its behavior locally. LIME generates these local explanations by perturbing ("shuffling") the feature values of the input observations to create a new dataset consisting of "synthetic" observations around an observation of interest. The responses of the complex model to these synthetic observations are then used to train a simpler, interpretable model, such as a linear regression or decision tree. This local surrogate model aims to capture how the original complex model behaves in the vicinity of the specified observation, providing insights into which features significantly influence the output and how they do so. LIME can be applied to any model type without any changes to the underlying algorithms.

LIME follows these steps:

1. Fit the complex model f
2. Choose an instance x of interest for which the prediction is to be explained
3. Transform the instance into a binary vector indicating the presence or absence of "interpretable components". Interpretable components are feature values, not features themselves.
4. Create a neighbourhood of n observations around x by perturbing interpretable components (sampling presence or absence of interpretable components randomly from $[0,1]$).
5. Use the fitted complex model f to create predictions for the generated neighbourhood of observations.
6. Weight the generated observations around x by a weight kernel π_g based on their distance to the instance x .
7. Sample a number of observations according to their weight.
8. Fit a local surrogate, interpretable model g to the sampled observations, using a set of k features, that minimizes the discrepancy \mathcal{L} between g and the complex original model f .



Source: Molnar Figure 9.5

Figure 16.10: Illustration of LIME

- The usual local surrogate model is a ridge regression model.
- The k features are identified by forward selection, by LASSO, or simply by selecting k features with the highest regression weights.

In principle, LIME aims to identify the explanation model as that local surrogate model g that minimizes the sum of the discrepancy between surrogate and original model plus a complexity penalty $\Omega(g)$ for the surrogate model:

$$\xi(x) = \underset{g \in G}{\operatorname{argmin}} \mathcal{L}(f, g, \pi_x) + \Omega(g)$$

However, in practice, as noted in the LIME procedure steps, the analyst must select the number of features and the type of local surrogate model. LIME only minimizes the discrepancy to the complex model; the complexity is determined by the analyst.

Figure 16.10 illustrates the principles behind LIME. The top left panel shows a classification problem with two predictors x_1 and x_2 and a somewhat complicated decision boundary.

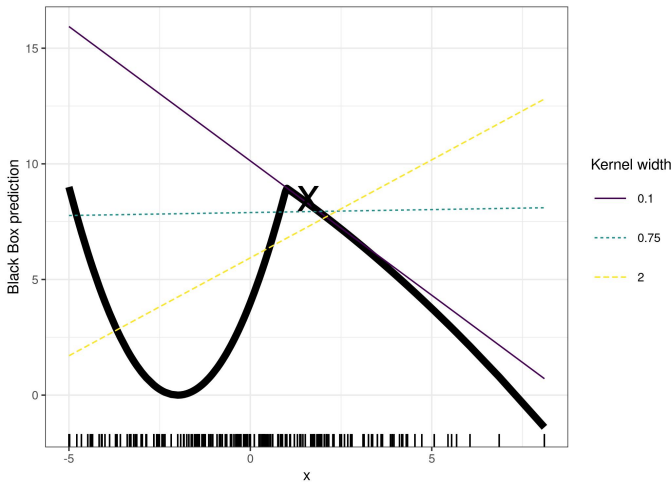


Figure 16.11: LIME results depend on choice of weight kernel

The top right panel highlights the target observation x in yellow and shows the observations created by perturbing the feature values as black dots.

The bottom left panel indicates the weighting of observations in the neighbourhood, bigger dots indicate higher weights. This weighting is induced by the distance function and the weight kernel. The distance function specifies how distances in the feature space are measured, for example Euclidean, Chebyshev or other metrics. The weight kernel uses the distances to assign weights. For example, a Gaussian kernel will assign weights according to a normal distribution density function, and an exponential kernel will assign weights according to the exponential of the distance, with higher weights for smaller distances and lower weights for larger distances.

The bottom right panel shows the class predictions of the simple surrogate model that has been fitted to the weighted samples. The white line shows the linear decision boundary of the surrogate model, the superimposed plus and minus signs the predictions of the surrogate model. The local surrogate model is quite accurate for values in the vicinity of the focal observation, but not very good for points far away from it.

Figure 16.11 illustrates the strong dependence of LIME results on the kernel, in particular the width of the kernel. The "X" marks the focal observation, and the graph plots the values predicted by the complex model against the values predicted by the simple model for different kernel widths. The point that Figure 16.11 makes is that not only the strength but also the sign or direction of the relationship between the predictions of the two models can change, depending on the choice of kernel.

The following Python code block uses LIME to provide an explanation for the predictions of a decision tree classifier.

```
import sklearn.tree
# Using a deep decision tree as black box model
dt = sklearn.tree.DecisionTreeClassifier(max_depth=8)
dt.fit(x, y)
```

The lime package provides the LimeTabularExplainer for regular datasets consisting of rows and columns. LIME can also be applied to other types of data such as images, see below.

```
import lime, lime.lime_tabular
# Create the explainer
explainer = lime.lime_tabular.LimeTabularExplainer(
    x.to_numpy(),
    feature_names=x.columns,
    discretize_continuous = True,
    mode='regression',
    verbose=True)
```

With the explaining model created, specific data instances can be explained using the explain_instance() method. The following example code explains the prediction for the 8th instance in the training data set that is generated by the predict() method of the regression tree object. The local neighbourhood is created with $n = 1000$ samples, the explanation is limited to $k = 5$ features, and the distance between observations for weighting them is determined by the Euclidean distance.

```
# Explain instance number 7
exp = explainer.explain_instance(
    x.to_numpy()[7],
    dt.predict,
    num_features=5,
    num_samples=1000,
    distance_metric='euclidean')
# Show weights as text and visualize
exp.as_list()
exp.as_pyplot_figure().show()
# Show complete explanation
exp.save_to_file('lime_explanation.html')
```

The results indicate the contribution of each feature value to the prediction. Note that continuous variables like temperature and windspeed have been discretized at various boundaries to form interpretable components. The visualizations in Figure 16.12 and Figure 16.13 shows those contributions as well.

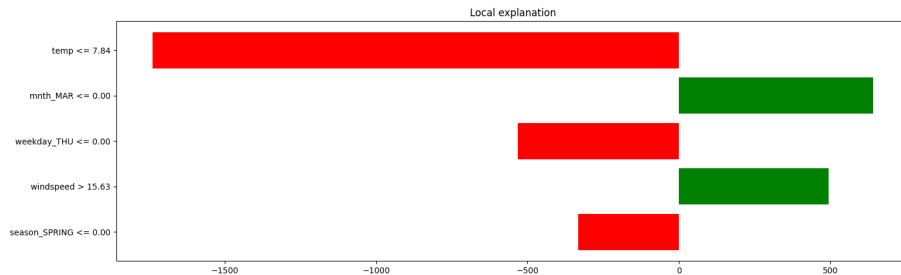


Figure 16.12: Weights of the LIME local surrogate model

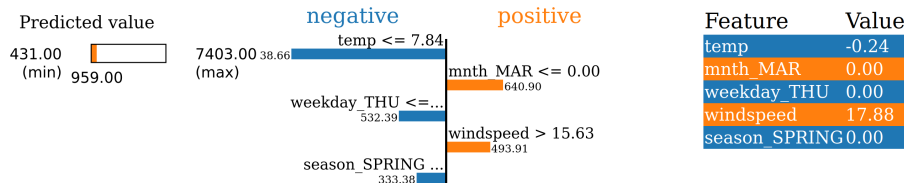
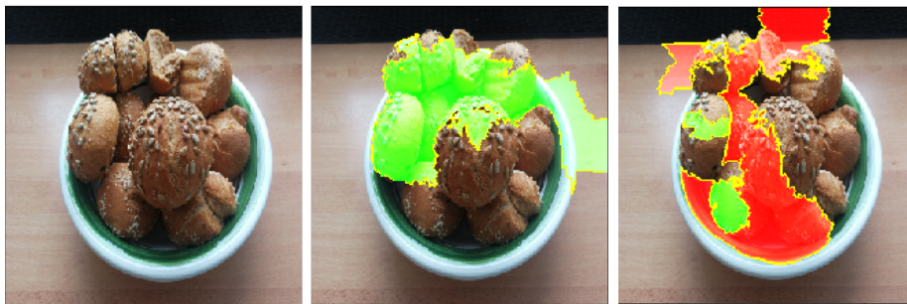


Figure 16.13: LIME results for explaining a specific instance

```
[ ('temp <= 7.84', -1738.6611673589232),
  ('mnth_MAR <= 0.00', 640.903016649792),
  ('weekday_THU <= 0.00', -532.3918204920352),
  ('windspeed > 15.63', 493.90993722141997),
  ('season_SPRING <= 0.00', -333.38496260796086) ]
```

Figure 16.13 consists of three parts. The left panel shows the prediction for this instance, 959 on the scale between the minimum (431) and maximum (7403). The right panel shows the actual feature values for the observation in table form. The center panel (which is identical to Figure 16.12) shows the negative and positive contribution of each interpretable component. For example, the contribution of $temp \leq 7.84$ is -1738.66 . This means that if the temperature were *not* equal to or below 7.84 degrees, the prediction would be 1738 higher than it is, that is, it would be 2697. Similarly, the feature $mnthMAR \leq 0.00$ contributes positively to the prediction (640.90). This means that if the month were March, the prediction would be 640.00 lower than it is.

LIME can also be used on other models such as image or text classification models. For example, Figure 16.14 shows the contributions of individual pixels of an image to the classification of that image. The left panel shows the original image, the middle panel the contributions for the label "bagel" and the right panel the contributions for the label "strawberries".



Molnar, Figure 9.8

Figure 16.14: LIME explanations for image classification

Further information on LIME, including many examples, can be found at the website for the Python package^a. More details on the theoretical foundation and principles of LIME can be found in the original paper^b.

^a<https://github.com/marcotcr/lime>

^b<https://arxiv.org/abs/1602.04938>

16.4.2 Shapley Additive eXplanations (SHAP)

Developed from the Shapley value – a concept from cooperative game theory originally used to determine the fair distribution of payoffs among players – SHAP applies this framework to the domain of machine learning, offering a theoretically sound approach to understanding model behavior. SHAP treats each feature value as a “player” in a game where the “payout” is the prediction itself. The Shapley value calculates the average contribution of each feature value across all possible combinations of feature values, providing a detailed view of how features interact and influence the model’s output. This approach not only yields a measure of global feature importance and local feature value importance, but also provides a way to understand the direction and magnitude of each feature’s impact on the prediction. SHAP is model-agnostic and can be applied to any machine learning model, from simple linear regressors to deep neural networks.

The main motivation in SHAP is the question of how much does a specific value x_j of feature j contribute to the overall prediction, when compared to the average prediction. As in LIME, the focus is on values of features, not the features themselves. In cooperative game theory, players cooperate in a coalition with other players and receive a certain profit from this cooperation. Shapley values are named after Lloyd Shapley who developed this concept in 1951 and received the Nobel prize in Economics for his work in 2012. Shapley values are a method for assigning fair payouts to the coalition’s players depending on their contribution to the total payout.

Formally, the Shapley value ϕ_i of player i is defined as:

$$\phi_i(v) = \frac{1}{n} \sum_{S \subseteq N \setminus \{i\}} \binom{n-1}{|S|}^{-1} [v(S \cup \{i\}) - v(S)]$$

where $v(\cdot)$ is the value function that describes the payout received in a game. The term $v(S \cup \{i\}) - v(S)$ is the marginal contribution of player i to a coalition of players S , that is, the difference in payout received by the coalition S plus the player i and the payout received only by the coalition S , without player i .

The binomial coefficient $\binom{n-1}{|S|}$ represents the number of possible ways to form a coalition of size $|S|$ of the set $N \setminus \{i\}$ of $n-1$ players (set N without player i). The marginal contribution is divided by this term.

The sum is taken over all all possible coalitions of the set N of players minus the player i . The sum is then divided by the number of players n .

Shapley values have some key theoretical properties that together ensure they describe a fair allocation of value or contribution to each player:

- *Efficiency*: The efficiency property states that all gains from cooperation are distributed among the players. Mathematically, this means the sum of the Shapley values for all players equals the total value (or payoff) that the coalition of all players achieves together. This ensures that the Shapley value captures the entire surplus generated by the group, leaving no residual value unallocated.
- *Symmetry*: The symmetry property ensures that if two players contribute equally to any coalition they are both members of, they receive the same proportion of the payoff. In other words, the Shapley value is identical for symmetric players, reflecting the principle of fairness in that contributions are rewarded equally without bias to factors unrelated to the contribution.
- *Additivity (or Linearity)*: Additivity is a property where the Shapley value of a game can be derived by summing the Shapley values of two separate games if a player's value in a combined game is the sum of their values in these separate games. This property allows for straightforward aggregation of separate contributions, simplifying the analysis and computation in more complex scenarios.
- *Dummy (or Null Player)*: A dummy (or null) player does not contribute to any coalition, i.e., the addition of this player does not change the value of any coalition. The dummy player property ensures that such a player receives a Shapley value of zero. This ensures that only contributors to a game's outcome are rewarded, maintaining the integrity of the allocation process.

In the context of interpretable machine learning, the players in a coalition are specific values of features (*not the feature themselves*). A coalition is a combination of different feature values. The presence of a feature value in a coalition means that the value of that feature is known and fixed. When a feature is not present in a coalition with a

specific value, its value is unknown. Determining the payout requires integrating or marginalizing over all values of those features $1 \dots p$ that are not in a coalition S .

$$v_x(S) = \int_{\mathbb{R}} \cdots \int_{\mathbb{R}} \hat{f}(x_1, \dots, x_p) d\mathbb{P}_{x \notin S} - E_x(\hat{f}(X))$$

This is clearly expensive to compute in practice and is therefore generally approximated by randomly permuting values (similar to how LIME constructs its local neighbourhood of observations) and then sampling the different possible coalitions from these permuted values.



Shapley Additive eXplanations (SHAP) is a Python package developed by authors of the paper that introduced Shapley values to interpretable machine learning¹. A thorough documentation, including an easy to read introduction and many examples are available², as well as the Python code for the implementation and multiple tutorial³. The remainder of this section illustrates the basic usage of SHAP.

This example uses a data set on California house prices that is part of the SHAP package. The target variable are house values in various districts, and predictors include median income, number of rooms, population, and similar variables. To illustrate the basic use of SHAP, this example fits a simple linear regression model to the data.

```
import sklearn
import shap
# Fit a simple regression model
X, y = shap.datasets.california(n_points=1000)
model = sklearn.linear_model.LinearRegression().fit(X, y)
```

The SHAP Explainer object is constructed using the fitted model's prediction function and the training data set. The explainer object can then be used to explain predictions in the same or another data set, e.g. just for a single new observation, or for the test or validation data set. The following Python code block illustrates this usage:

```
# Create the Explainer object:
explainer = shap.Explainer(model.predict, X)
# Compute the SHAP values:
shap_values = explainer(X)
```

The SHAP package provides different ways to visualize individual predictions. The *barplot* in Figure 16.15 shows the importance of feature values (*not the importance of*

¹<https://arxiv.org/abs/1705.07874>

²<https://shap.readthedocs.io/en/latest/index.html>

³<https://github.com/shap/shap>

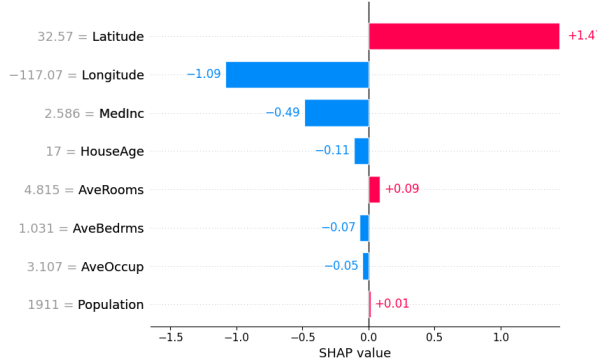


Figure 16.15: SHAP barplot for an individual prediction

features!) for an individual prediction, generated from the running example using the following Python code:

```
shap.plots.bar(shap_values[20])
```

For example, the fact that the house age is 17 contributes negatively to the prediction, whereas the fact that the average number of rooms of houses in the district is 4.815 contributes positively.

By averaging over all instances and their feature values, SHAP values can be aggregated to show the importance of the features themselves, shown in the bar plot in Figure 16.16, generated from the running example using the following Python code:

```
shap.plots.bar(shap_values)
```

The graph shows that latitude and longitude are the most important features, that is, their values on average contribute positively to the prediction.

Waterfall plots such as the one in Figure 16.17 explain how feature values combine to produce an individual prediction. In that figure, latitude being 32.57 contributes positively to the house price prediction, whereas longitude being -117.07 contributes negatively, etc. The total sum of contributions is the expected house price, shown at the very bottom of the waterfall plot. The waterfall plot was generated by the following Python code:

```
shap.plots.waterfall(shap_values[20])
```

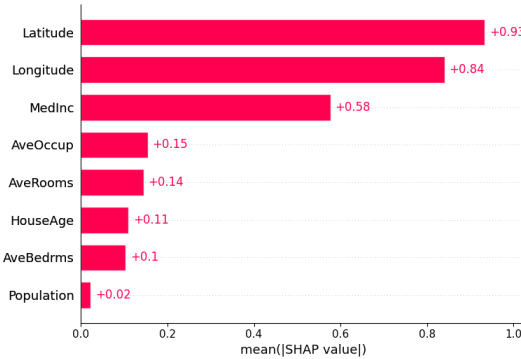


Figure 16.16: SHAP barplot of mean SHAP values for feature importance

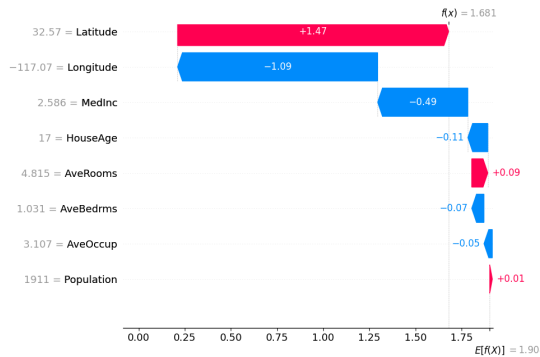


Figure 16.17: SHAP waterfall plot for an individual prediction

Beeswarm plots, such as the one in Figure 16.18 explain all feature values for all instances (represented by a dot). Each dot is particular feature value that occurs in the data. Dots are colour-coded to indicate whether that feature value makes a positive or negative contribution. Figure 16.18 shows that the median district income feature has a lot of feature values in the data set that contribute negatively, and a few values that make a highly positive contribution. Figure 16.18 was generated using the following Python code.

```
shap.plots.beeswarm(shap_values)
```

The *heatmap plot* in Figure 16.19 shows the SHAP values of feature values for all instances, and shows model prediction and global feature importance in the top and right rugs of the figure. Observations are ordered left to right. The bar plot in the right rug is the same as the feature importance bar plot in Figure 16.16. The figure shows

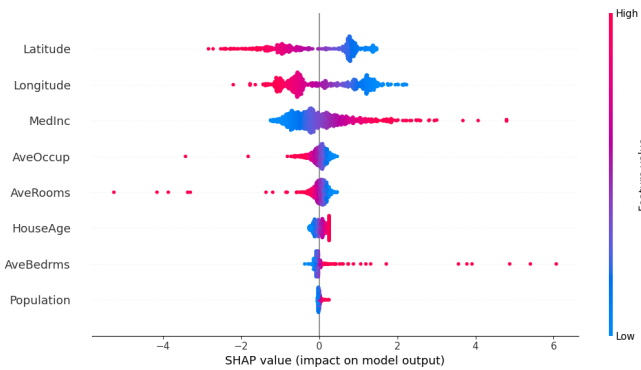


Figure 16.18: SHAP beeswarm plot

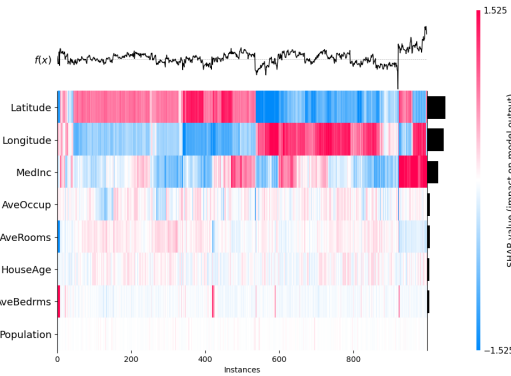
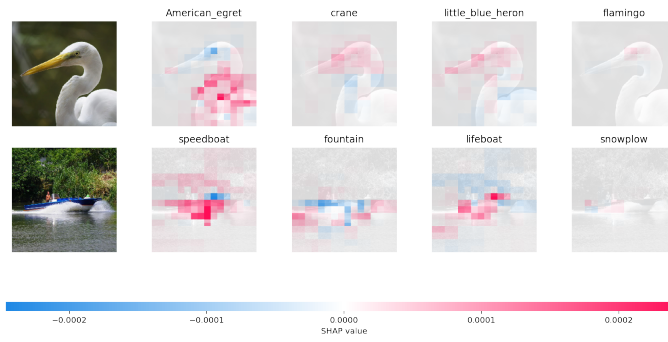


Figure 16.19: SHAP heatmap plot

clearly how some values of median district income contribute highly positively to predicted house prices, whereas some values of latitude appear to contribute negatively the predicted house price. Figure 16.19 was generated using the following Python code:

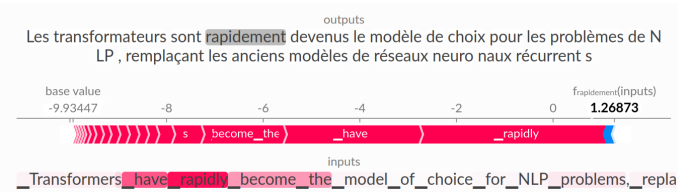
```
shap.plots.heatmap(shap_values)
```

As noted earlier, SHAP is model agnostic and can be applied to problems such as image classification and text classification as well. The SHAP package provides intuitive visualizations for the SHAP values in these cases. For example, Figure 16.20 shows the contribution of different image elements (groups of pixels) to classification probabilities for different target classes. In this case, the absence of a feature value is created by masking a group of pixels. Consider the top image in Figure 16.20. The overlaid SHAP values show that it is mostly the body or neck shape of the bird that contributes positively to a classification as an American egret, while the eye contributes negatively.



Source: <https://github.com/shap> (MIT License)

Figure 16.20: SHAP for image classification



Source: https://shap.readthedocs.io/en/latest/text_examples.html (MIT License)

Figure 16.21: SHAP for text classification

Figure 16.21 shows SHAP applied to text classification. Each token/word in a text is a feature that contributes to class membership probabilities.

16.5 Review Questions

Introduction to Interpretable Machine Learning

1. List at least three reasons why interpretable machine learning is crucial in modern AI applications.
2. What does it mean for a machine learning model to be "interpretable"? Give examples of features that contribute to a model's interpretability.
3. Discuss the trade-offs between model complexity and interpretability. Why might a more complex model be chosen over a simpler, more interpretable model?
4. Provide examples of how interpretable machine learning can lead to innovations and improvements in AI systems.
5. Why is the understanding of automated decisions particularly crucial in sectors like healthcare and finance?
6. Discuss the relationship between interpretable machine learning and the ethical responsibilities of AI developers.
7. Discuss the role of interpretable machine learning in ensuring safety, compliance, and reliability of AI systems.

8. What is the importance of interpretable machine learning in detecting biases and ensuring fairness in model predictions?
9. What factors should be considered when deciding between intrinsic and post-hoc interpretability methods in a given application?
10. Compare and contrast local and global interpretation methods in terms of their applications and benefits.
11. How do local and global interpretation methods complement each other in providing a comprehensive understanding of machine learning models?
12. How do intrinsic and post-hoc methods affect the user's trust and acceptance of machine learning models?

Intrinsically Interpretable Models

13. How do t-values contribute to understanding the relative importance of features in a linear regression model?
14. What are decision trees, and how are they used in both regression and classification contexts in machine learning?
15. Describe the structure of a typical decision tree and explain how decisions are represented within this structure.
16. Explain why decision trees do not require normalization or scaling of data before analysis. What advantages does this present in terms of data preprocessing?
17. Discuss the advantages of using decision trees in terms of algorithmic transparency. How does this feature contribute to their interpretability?
18. Explain how decision trees handle non-linear relationships and the significance of this capability.
19. Discuss how decision trees manage to capture interactions between variables without explicit programming for these interactions.
20. What is overfitting in the context of decision trees, and why are decision trees particularly susceptible to it?
21. Explain the process of making binary splits in decision trees, particularly how regions and leaf nodes are determined.
22. Explain the various stopping criteria used in the construction of decision trees. Compare them in terms of their impact on tree complexity and performance.
23. Discuss the implications of high variance in decision trees. What techniques can be used to mitigate this issue?

Global Model-Agnostic Methods

24. What are Partial Dependence Plots (PDPs) and what do they show in a machine learning model?
25. Explain the assumption of feature independence in PDPs and its implications.
26. Describe a scenario where the assumption of feature independence in PDPs might lead to incorrect conclusions about feature importance.
27. How can ICE plots help in identifying outlier effects or heterogeneity in data predictions?

28. Describe how ALE plots differ from PDPs in their approach to understanding feature effects.
29. Explain the computation process of ALE plots and how it differs fundamentally from PDPs in handling data within local windows.
30. Discuss how ALE plots handle correlated features and why this is beneficial.
31. What is Permutation Feature Importance and how is it calculated?
32. Explain global surrogate models in interpretable machine learning.
33. Discuss the potential risks or drawbacks of using global surrogate models as a tool for interpretability. What should practitioners be wary of?
34. Compare and discuss the strengths and weaknesses of the global model-agnostic methods covered (PDP, ICE, ALE, PFI, and global surrogate models).

Local Interpretable Model-agnostic Explanations (LIME)

35. What are local model-agnostic interpretative methods, and how do they differ from global interpretative methods?
36. Outline the step-by-step process of generating a LIME explanation for a given data instance.
37. Explain the purpose of transforming an instance into a binary vector in the LIME process. What does this achieve in terms of interpretability?
38. Discuss the role of perturbing input data points in the LIME process. What is the purpose of creating "synthetic" samples?
39. How does LIME ensure that the local surrogate model accurately reflects the behavior of the complex model in the vicinity of a specified data point?
40. Identify potential pitfalls when applying LIME to a dataset with highly correlated features. How does this correlation affect the integrity of the explanations provided?
41. Provide a critical analysis of how LIME handles cases where the local decision boundary is highly non-linear. What are the challenges and how might LIME overcome them?
42. Reflect on the dependency of LIME's explanations on the kernel function. How can different kernels alter the interpretation of feature contributions?

Shapley Additive eXplanations (SHAP)

43. Explain the concept of Shapley values in the context of cooperative game theory.
44. Discuss the significance of treating each feature as a "player" in the context of SHAP. How does this perspective aid in understanding model predictions?
45. Describe the four key theoretical properties of the Shapley value and explain how each contributes to fairness and accuracy in SHAP.
46. Outline the steps to compute SHAP values for a machine learning model.
47. Describe how SHAP values are visualized and interpret such a visualization.
48. What insights can be gained from SHAP waterfall and beeswarm plots?
49. How can SHAP be integrated into the model development process to improve model design and feature engineering?
50. Discuss the potential for SHAP to be used in regulatory compliance, specifically

in industries like finance and healthcare. What advantages does SHAP offer in explaining model decisions to regulators?

Chapter 17

Analytics at Industrial Scale

Sources and Further Reading

The material in this chapter is based on the following sources.

Tom White (2012) *Hadoop – The Definitive Guide*. 3rd edition. O'Reilly Media. Sebastopol, California, US.

This book introduces in great detail the main concepts of Apache Hadoop and provides step-by-step introduction to setting up a Hadoop cluster, using the Map-Reduce API, submitting and managing jobs. The book also provides a chapter on related projects, such as Hive and Pig. It was written for Hadoop 2.3, which introduced the Map-Reduce 2.0 API and YARN. While older, it can still serve as a comprehensive reference also to more recent Hadoop versions.

Hrishikesh V. Karambelkar (2018) *Apache Hadoop 3 Quick Start Guide*. Packt Publishing. Birmingham, UK.

This quick start guide is more recent and written for Hadoop 3.0. It provides only a brief overview of the Hadoop system, focusing on cluster installation, configuration, and management, rather than Map-Reduce or data management.

Bill Chambers and Matei Zaharia (2018) *Spark – The Definitive Guide*. O'Reilly Media. Sebastopol, California, US.

This is a very comprehensive book, written for Spark 2.0. Matei Zaharia is the original designer and lead developer of the Spark project. The book covers all aspects of Spark. All examples are provided both in Scala and in Python. While a little older at this point,

it is very instructive and easy to follow with many practical points of how to get the most out of Spark.

Jules Damji et al. (2020) *Learning Spark – Lightning-Fast Data Analytics*. 2nd edition. O'Reilly Media. Sebastopol, California, US.

This book, while not quite as comprehensive as the one by Chambers & Zaharia, is newer and targets Spark 3.0. It also covers all aspects of Spark, from installation and configuration, to programming the SQL, streaming, and machine learning components of Spark. The book provides most examples in both Scala and Python.

Resources

Complete implementations of all examples in this chapter are available in the following GitHub repos.

For the Hadoop MapReduce examples:

<https://github.com/jevermann/busi4720-hadoop>

The project can be cloned from this URL:

<https://github.com/jevermann/busi4720-hadoop.git>

For the Spark examples:

<https://github.com/jevermann/busi4720-pyspark>

The project can be cloned from this URL:

<https://github.com/jevermann/busi4720-pyspark.git>

17.1 Introduction

Big data analytics refers to the process of examining large and varied data sets – *big data* – to uncover hidden patterns, unknown correlations, market trends, customer preferences, and other useful information that can help organizations make better-informed business decisions. Driven by specialized analytics systems and software, big data analytics can lead to more effective marketing, new revenue opportunities, better customer service, improved operational efficiency, competitive advantages over rival organizations, and other business benefits.

The term "big data" is typically associated with three key concepts: data volume, data variety, and data velocity. Together, these three Vs define the challenges and opportunities that organizations face when managing and analyzing massive data sets. The emergence of the fourth and fifth Vs, data veracity and data value, reflect the increasing need to ensure the reliability of data and the importance of deriving meaningful insight

from it.

Volume represents the quantity of data that is generated and stored. It is one of the primary characteristics of big data. Organizations collect data from a variety of sources, including business transactions, smart (IoT, Internet-of-Things) devices, industrial equipment, videos, social media, and more. As data volume increases, the value and potential insights also increase, but it requires more effective data management and processing technologies. The challenge lies not only in the storage but also in dynamically scaling resources to process this vast amount of data efficiently.

Variety refers to the different types of data that are available, both structured and unstructured. Traditional data types were structured and fit neatly in a relational database. However, with the advent of big data, data comes in new unstructured forms such as text, images, videos, and social media posts. The variety in data types frequently requires additional preprocessing to derive meaning and supports analytics that can lead to enhanced business insights.

Velocity is the speed at which the data is generated, collected, and processed. High velocity of big data comes from connected sensors, smart devices, web page streaming, and other live feeds. Managing the velocity of data entails not only processing these streams effectively but also ensuring that data analysis and decision making are conducted in a timely manner.

Veracity refers to the accuracy of data. Big data veracity deals with the assurance of quality and accuracy of data considering its volume, variety, and velocity. This characteristic is crucial as it affects every decision made based on big data. Poor data quality can lead to incorrect conclusions. Therefore, organizations must invest in analytic processes that validate and cleanse data to ensure accuracy and usefulness.

Value value is about turning big data into business value. This is the most important V of all, as it determines whether the data collected and analyzed are of any use to the organization. The main challenge is to sift through vast lakes of data and find what is relevant and can be turned into actionable insights that can lead to cost reductions, improved efficiency, or new revenue opportunities.

An example of a big data use case is the Conseil Europeen pour la Recherche Nucléaire (CERN). CERN is the world's largest physics laboratory, an international cooperation that researches high-energy particle physics located in Switzerland and France. When physics experiments are running, the CERN data center processes about one petabyte (one million gigabyte) of data per day¹. Preparations are under way for new physics experiments that require storage and analysis of 600 petabytes of data. Table 17.1 provides an overview over the CERN data centre infrastructure to manage this volume and this velocity of data.

¹<https://www.home.cern/science/computing/storage>, last accessed June 10, 2024

Servers	≈ 12000
CPU Cores	≈ 350000
Disks	≈ 220000
Total Disk Space	≈ 950000 TB
DB Transactions per second	≈ 20000
File Transfer Throughput	≈ 100 Gb/s

Table 17.1: Data management infrastructure at CERN²

Source: CERN.org with permission

Figure 17.1: CERN data centre images

Figure 17.1 shows images of the CERN data centre infrastructure. Racks are filled with multiple computer servers, each with their own attached data storage, and multiple racks are lined up in rows. Computers are networked, first within a rack, then between racks, and the data center is connected to the larger internet. Network connections are fastest between computers within the same rack, then between computers in different racks, and are slowest across the internet.

Clearly, the volume of data being stored, and the velocity with which it is generated requires analytics techniques and software tools that go beyond those of single-machine or desktop solutions examined so far, like R, Numpy or Pandas, and that are suitable to efficiently use infrastructure like the one shown in Figure 17.1. The following sections introduce big data analytics, or data analytics at industrial scales.

17.2 Hadoop



Apache Hadoop is an open-source system for distributed data storage and distribute computation on data. Initially released in 2006, it is maintained by the Apache Foundation. It was originally inspired by the

¹<https://commons.wikimedia.org>.

²<https://www.cern.ch/en/science/computing/data-centre>, last accessed June 10, 2024 `Apache_Hadoop.png`

Google File System (GFS) that Google used to manage its web search data³, and the Google MapReduce technique for computations on distributed data⁴. Early successful uses cases at Yahoo in 2009 and at Facebook in 2012 drove industry adoption and further research interest. Hadoop relies on a distributed file system called HDFS ("Hadoop Distributed File System") that distributes data storage across a cluster of computers. In contrast to other solutions at the time, the clusters did not require specialized hardware or software, which made the system very attractive to organizations.

The main principle of data processing with Hadoop is *data locality*: Instead of moving data across network connections to the computer where computation takes place, which is expensive and slow, computation is moved to where the data is stored. In other words, data is processed where it is stored, in a distributed fashion.

The benefits that make Hadoop attractive are its reliability, scalability, cost effectiveness and cloud support.

Reliability Hadoop is highly reliable. It stores multiple copies of data across different computer nodes in a network, ensuring that the system can tolerate the failure of any network node without data loss. This redundancy provides fault tolerance, as data is automatically re-replicated from the remaining copies when a network node fails, allowing data processing to continue uninterrupted.

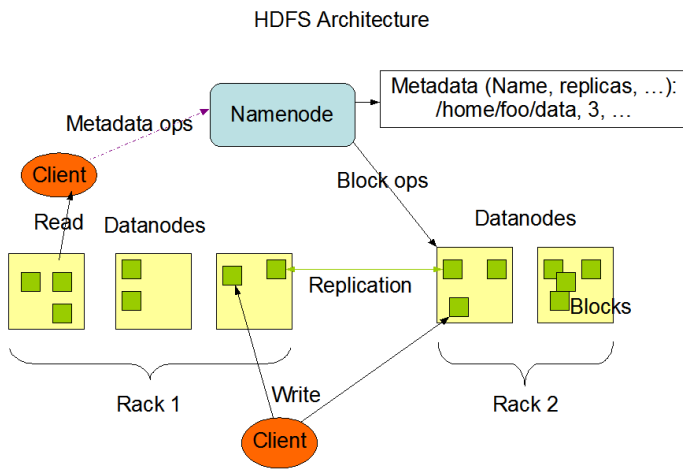
Scalability Scalability is one of Hadoop's core strengths. It can store and distribute very large data sets across hundreds to thousands of inexpensive servers that operate in parallel. This distributed computing model allows businesses to scale up or down efficiently without downtime. As more servers are added, Hadoop continues to increase its storage capacity, processing power, and throughput performance proportionally.

Cost Effectiveness Hadoop is cost-effective. As open-source software, combined with the use of commodity hardware, it significantly reduces the cost of a system capable of storing and processing enormous amounts of data. The cost savings are not just in terms of hardware but also in scalability. Businesses can start with what they need and increase their system size as they grow while maintaining a low cost per terabyte. Hadoop 3 provides support clusters with more than 10,000 computers.

Cloud Support Finally, Hadoop supports cloud-based services, which allows businesses to deploy Hadoop clusters in cloud environments. This capability means that organizations can benefit from cloud computing features, such as massive scalability, on-demand resource allocation, and utility-based cost structures. Most major cloud

³Ghemawat, S., Gobioff, H., & Leung, S. T. (2003, October). The Google file system. In Proceedings of the nineteenth ACM symposium on Operating systems principles (pp. 29-43).

⁴Dean, J., & Ghemawat, S. (2008). MapReduce: simplified data processing on large clusters. Communications of the ACM, 51(1), 107-113.



Source: Apache Foundation (<https://hadoop.apache.org/docs/>)

Figure 17.2: HDFS architecture

vendors, such as Amazon Web Services (AWS), Google Cloud Platform (GCP), and Microsoft Azure offer Hadoop or equivalent services to their customers.

Hadoop consists of three main components. HDFS is the storage layer of Hadoop. It is a distributed file system designed to run on commodity hardware. MapReduce is a programming model and implementation for processing large data sets with a parallel, distributed algorithm, and YARN is the resource management layer of Hadoop.

17.2.1 HDFS

HDFS is designed and optimized for storing very large datasets, ranging from a few gigabytes to hundreds of terabytes in size. HDFS is designed based on a number of fundamental principles: Data in HDFS is written and read linearly, and must be processed one item at a time. For analytics applications, this means that applications cannot read arbitrary data or move back and forth in the data set. Once a file is written and closed, it cannot be changed, other than by appending data or truncating the file. This means that existing data in the file cannot be overwritten. If necessary, a new file must be created. HDFS is built around the idea of streaming data access, that means it emphasizes high throughput over low latency. In other words, accessing the same amount of data from one continuous file is much faster than accessing the same amount of data from multiple separate files.

Figure 17.2 shows the architecture of HDFS. An HDFS cluster consists of a single Namenode (optionally a secondary/backup NameNode for fault tolerance), which is a server that runs software to manage the file system namespace and regulates access to files by client applications. The Namenode keeps the directory of all files in the file system, and tracks where on the cluster the file data is kept. The Namenode

does not store actual data. Instead, it stores metadata, such as the location of blocks stored on the Datanodes, access permissions, and other information. It provides file operations such as opening, closing, renaming of files. Client applications communicate with the Namenode whenever they wish to locate a file, or when they want to add/copy/move/delete/... a file. The Namenode responds to the requests by returning a list of relevant DataNode servers where the data lives.

Datanodes manage the storage attached to the computers they run on and serve read/write requests from the file system's client applications. HDFS splits large files into blocks (the default block size is 128MB) and distributes them across nodes in a cluster to enable high throughput access to data. The file system also replicates each data block multiple times (by default 3 times) across different nodes to ensure reliability and fault tolerance. Datanodes store and retrieve blocks when they are told to (by client applications or the NameNode), and they report back to the Namenode periodically with a list of blocks that they are storing.

Working with HDFS is very similar to working with the local file system on a desktop machine. In other words, the cluster is transparent. HDFS file commands are based on the standard Unix/Linux file commands to manipulate files and directories. The set of basic HDFS commands is shown in Table 17.2. Because HDFS emphasizes throughput over latency, file operations that only involve small amounts of data, tend to be much slower than on a local file system.

In addition to the command line interface, the Hadoop Namenode also typically provides web-based access. For example, the Namenode overview can be accessed at <http://localhost:9870> (substitute another URL or host name if necessary) and will show the status of the datanodes in the cluster as well. This also allows browsing the file system and performing file operations, including uploading and downloading of files from and to the local filesystem through the HDFS explorer at <http://localhost:9870/explorer.html#/> (substitute another host name if necessary). Figure 17.3 shows a screenshot of the HDFS explorer.

The following exercises illustrate the use of the HDFS. Use the `hdfs dfs` command to interact with the distributed file system. The commands are similar to the regular Linux commands to interact with files.

On a single-machine cluster, the Hadoop Namenode, Datanode, and the YARN software applications are all installed on the same computer. Start the Hadoop cluster Namenode, Datanode, and YARN service as follows:

```
sudo systemctl start hadoop.service
```

Download an event log file to your local file system and then put the event log onto the Hadoop Distributed File System:

```
wget https://evermann.ca/busi4720/eventlog.short.log
hdfs dfs -put eventlog.short.log
```

The screenshot shows a web-based HDFS browser interface. At the top, there's a header bar with tabs for 'Hadoop', 'Overview', 'Datanodes', 'Datanode Volume Failures', 'Snapshot', 'Startup Progress', and 'Utilities'. Below the header is a search bar and a toolbar with icons for file operations. The main area is titled 'Browse Directory' and shows a list of files and directories under '/user/busi4720'. The list includes:

Name	Size	Last Modified	Replication	Block Size
.sparkStaging	0 B	Apr 03 15:56	0	0 B
commits	0 B	Apr 03 15:50	0	0 B
eventlog.short.log	6.66 MB	Apr 03 15:42	1	128 MB
hamlet	0 B	Apr 03 15:56	0	0 B
hamlet.out	0 B	Apr 03 15:59	0	0 B
hamlet.txt	223.19 KB	Apr 03 16:16	1	128 MB
metadata	45 B	Apr 03 12:49	1	128 MB
mushrooms.csv	4.21 MB	Apr 03 12:33	1	128 MB
offsets	0 B	Apr 03 16:49	0	0 B
state	0 B	Apr 03 16:46	0	0 B

Figure 17.3: HDFS Explorer

hdfs dfs -cat	Print a file to standard output
hdfs dfs -cp	Copy a file or directory
hdfs dfs -df	Display free space
hdfs dfs -du	Display disk usage
hdfs dfs -get	Copy files to the local file system
hdfs dfs -head	Print the first kilobyte of a file
hdfs dfs -ls	List files and directories
hdfs dfs -mkdir	Make a directory
hdfs dfs -mv	Move a file or directory
hdfs dfs -put	Copy files from the local file system
hdfs dfs -rm	Remove files or directories
hdfs dfs -rmdir	Removes a directory
hdfs dfs -tail	Print the last kilobyte of a file
hdfs dfs -concat	Concatenate existing files into a target file

Table 17.2: Basic HDFS file system commands

Display the start and end of the from the HDFS:

```
hdfs dfs -head eventlog.short.log
hdfs dfs -tail eventlog.short.log
```

Show disk usage and disk free space:

```
hdfs dfs -du  
hdfs dfs -df
```

Copy the event log:

```
hdfs dfs -cp eventlog.short.log eventlog.copy.log
```

List all files:

```
hdfs dfs -ls
```

Hands-On Exercise

Exercise 1: Accessing HDFS

1. View the list of files on the HDFS file system.
2. Create a new directory in the HDFS named "testdir".
3. Verify that 'testdir' has been created by listing all files.

Exercise 2: Manipulating Files in HDFS

1. Use a text editor to create a text file called "example.txt" on your local file system and write "Hello HDFS" into it.
2. Copy "example.txt" from your local file system to the directory "testdir" in the HDFS.
3. Read the contents of the file from the HDFS.
4. Delete "example.txt" from the HDFS.

Exercise 3: Understanding HDFS Block Size

1. Create a large text file (e.g., larger than 128MB, the default block size in many HDFS installations) named "largefile.txt" on your local file system.
2. Copy "largefile.txt" to HDFS:
3. Use the `hdfs fsck` command with the `-files -blocks` options to check how HDFS has stored the file in terms of blocks.
4. Observe and note the number of blocks the file is split into and their locations.

17.2.2 Map-Reduce

MapReduce is a programming model and a software implementation for processing large data sets with a parallel, distributed algorithm. The main idea is to use the distributed data storage of HDFS also for computation, that is, computation is performed

on the nodes where the data is stored. This means that each block of a file or data set is initially processed independently of all other blocks.

MapReduce is designed with two main functions: *Map* and *Reduce*. Blocks of a file are independently processed by the *Map* function in a completely parallel manner, each on the computer where it is stored. The MapReduce framework then sorts the outputs of all the *Map* functions, which are then input to the *Reduce* function. Typically both the input and the output of the MapReduce application are stored in HDFS. MapReduce allows for massive scalability across hundreds or thousands of servers in a Hadoop cluster.

Not only are input data and results stored on HDFS but intermediate results (between *Map* and *Reduce*) are stored on HDFS as well. For some applications, the size of this intermediate data may be larger than that of the input data. The lack of in-memory storage or data streams means that the performance of MapReduce is sometimes limited by disk performance. Moreover, because data must be read and processed linearly, the *Map* and *Reduce* functions are necessarily stateless, that is, they are limited in how much of a "memory" they can maintain. Finally, MapReduce supports only non-iterative, acyclic data flows. While multiple *Map–Reduce* phases can be executed in sequence, it is not possible for data to be processed multiple times, in a cycle.

The basic steps of MapReduce are the following:

1. Map

- The *Map* function reads key–value pairs of input data⁵
- For each input key and value, the *Map* function outputs a list of key–value pairs:

$$\textit{Map} : (key1, value1) \rightarrow list(key2, value2)$$

- Multiple instances of *Map* operate in parallel, one on each block of data

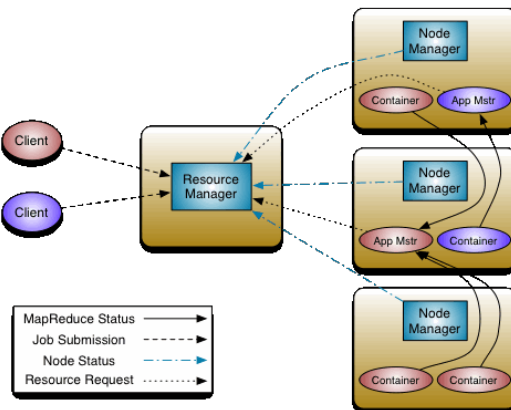
2. Shuffle

- The shuffle stage distributes the output of the *Map* function based on keys produced by *Map*
- All values for the same key are sent to the same instance of the *Reduce* function.

3. Reduce

- The *Reduce* function processes all values for a given key
- There are as many instances of *Reduce* as there are unique values for input key *key2*

⁵By default, for text input, each line is a key–value pair, separated by the first tab character



https://hadoop.apache.org/docs/current/hadoop-yarn/hadoop-yarn-site/yarn_architecture.gif

Figure 17.4: Executing MapReduce job on YARN cluster manager

- For each input key and its values, the Reduce function outputs a list of key–value pairs

$$\text{Reduce} : (key2, list(value2)) \rightarrow list(key3, value3)$$

YARN is the resource management layer of Hadoop. YARN consists of a central Resource Manager, which manages the use of resources across the cluster, and NodeManager agents, which monitor the processing of operations on individual cluster nodes.

Running a MapReduce application on a Hadoop cluster involves submitting the application to the YARN resource manager. The YARN resource manager creates an application master process for each MapReduce application that is submitted to it. The application master process tracks the status of the tasks, that is, the Map and Reduce instances, within an application. The application master requests resources from the resource manager and then distributes task to nodes in the cluster. The node managers manage the local resources and report their status to the resource manager. They execute the tasks assigned by the application master in containers on each node. Figure 17.4 shows the interplay between these components.

For a typical MapReduce application, the user specifies an input directory, possibly with multiple data files, to be processed. Recall that data files are distributed in blocks across the cluster node. A separate Map instance is executed for each input data block, on the node where the input data block is located. This means that any necessary program files are sent to a node as required. The shuffle phase sorts the output by key and moves data with the same key to the same node. While every instance of Reduce "sees" all values for the same key, a Reduce instance may process the values for multiple keys, depending on the number of unique key values and the number of nodes available in the cluster.

Example

While Hadoop MapReduce is natively programmed in Java, Hadoop Streaming allows Map and Reduce functions as executable programs, for example using Python. This example shows the basic operation of the Map and Reduce functions. The aim is to count the number of distinct words in a large text document. Both the Map and Reduce functions are implemented in Python.

The Map function in the following Python code block reads lines from standard input. Hadoop Streaming is responsible for feeding those lines from the specified input to the Map function. After whitespace is stripped from the beginning and end of each line, the line is split into words. For each word, the Map outputs a key–value pair, separated by a tab character. The key is the word, the value is the number 1.

```
#!/usr/bin/env python
import sys

for line in sys.stdin:
    line = line.strip()
    words = line.split()
    for word in words:
        print ('{}\\t{}'.format(word, 1))
```

Every instance of the Reduce function sees all values for a key, but may be processing the multiple keys. The following Reduce implementation reads key–value pairs from the standard input. It splits them on the tab character and maintains a dictionary of word counts. For each new key–value pair it increments its counter by the value it has just read. Finally, it outputs its results in the form of key–value pairs.

```
#!/usr/bin/env python
import sys

word_counts = dict()

for line in sys.stdin:
    word, count = line.split('\\t', 1)
    count = int(count)

    if word not in word_counts:
        word_counts[word] = count
    else:
        word_counts[word] = word_counts[word] + count

for word, count in word_counts.items():
    print ('{}\\t{}'.format(word, count))
```

Before submitting this application as a job to a Hadoop cluster, it is useful to run it on the local machine and local filesystem. Download the program files and a data file and make the downloaded files executable:

```
wget https://evermann.ca/busi4720/map.py
wget https://evermann.ca/busi4720/reduce.py
wget https://evermann.ca/busi4720/hamlet.txt
chmod +x *.py
```

Then, run the Map function and view its output to understand how it works. The “|” symbol in the following bash code is a *pipe* that pipes the output of one command into the next command. Here, the output of `cat`, which simply shows the contents of a file, are piped into the map function defined above, and its output is redirected with the `>` symbol to a file. The `less` command shows the contents of that file one line in an interactive way.

```
cat hamlet.txt | ./map.py > map.out
less map.out
```

Next, run the Reduce function and view its output. The `sort` command sorts the content of text files. The `k2` option tells it that the sort key is the second column (word count) in the file, the `rn` option is to reverse the sort order (highest word count first).

```
cat map.out | ./reduce.py > reduce.out
sort -k2 -rn reduce.out | less
```

Once the Map and Reduce functions work as expected on the local file system, they can be run on the Hadoop cluster. To do this, first put the input file(s) into their own directory on the HDFS:

```
hdfs dfs -mkdir hamlet
hdfs dfs -put hamlet.txt hamlet
hdfs dfs -ls hamlet
```

Next, run the MapReduce application on the Hadoop cluster using Hadoop Streaming. The command arguments are self-explanatory, but note the use of the `file` arguments to tell Hadoop to move the program code to the nodes where the data are located. This illustrates the fundamental Hadoop principle that computation is moved to the data, not the other way around.

```
mapred streaming \
-input hamlet -output hamlet.out \
-mapper map.py -reducer reduce.py \
-file map.py -file reduce.py
```

The Hadoop cluster will now process the application, provide statistics and information about it as it is executed, and then report successful execution. The output directory, hamlet.out contains one result file from each instance of Reduce.

Download the results to the local file system and sort and view them:

```
hdfs dfs -ls hamlet.out
hdfs dfs -get hamlet.out/part-
cat part-* | sort -k2 -rn | less
```

Case Study



Source: IEEE

This case study illustrates the scalability of Hadoop in real business use case⁶. One aspect of business process mining is the discovery of process models from event logs. Event logs are generated by process-aware information systems and can grow to significant size. Event logs may be stored in a distributed way, either on those systems where they are generated, or in a dedicated business analytics Hadoop cluster. This suggests that a distributed implementation of process discovery algorithms using the MapReduce framework could provide significant scalability and performance advantages.

The α miner and the Flexible Heuristics Minder (FHM) were implemented in MapReduce, the α miner requiring 2 Map–Reduce phases, while the FHM required 5 Map–Reduce phases. Experiments were conducted on a randomly generated process with 47 different types of activities. From this process, 5 million traces were simulated, yielding an event log of 80GB. To examine the effect of cluster size on performance, a single node cluster with 2 CPUs was chosen as a baseline and compared to a 10-node cluster with 2 CPUs on each node, and a 10-node cluster with 10 CPUs on each node.

While the above word count example used simple data types for the keys and values, the data can be arbitrarily complex, as long as a function to compare keys is provided so that the shuffle phase can decide when two keys are the same and send values to the same Reduce instance. For example, the MapReduce implementation of the FHM used tuples of two elements for some keys and sets of multiple elements for some values:

```

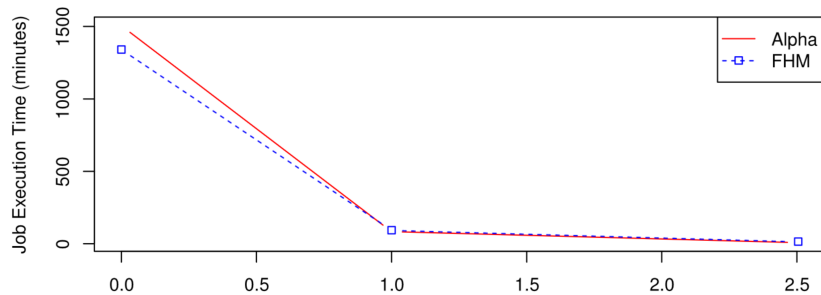
map1:(Int,Text) → set(CaseID,(Event,TimeStamp))
shuffle1:set(caseID,(Event,TimeStamp)) → (CaseID, set(Event,TimeStamp))
reduce1:(CaseID, set(Event,TimeStamp)) → set((Event,Event),(Int,Bool,Int))
combine2:set((Event,Event),(Int,Bool,Int)) → set((Event,Event),(Int,Bool,Int))
reduce2:((Event,Event),set(Int,Bool,Int)) → set(c,(Event,Event,Int,Float))
reduce3:set(c,(Event,Event),set(Int,Float)) → set(c,(Event,Event))

map4:(Int,Text) → set(CaseID,(Event,TimeStamp))
shuffle4:set(CaseID,(Event,TimeStamp)) → (CaseID, set(Event,TimeStamp))
reduce4:(CaseID, set(Event,TimeStamp)) → set((Event, set(Event),Bool),Int)
reduce5:((Event, set(Event),Bool),set(Int)) → ((Event, set(Event),Bool),Int)

```

The experimental results, shown in Figure 17.5 and Table 17.3, show the performance advantages delivered by parallel processing using multiple compute nodes with multiple CPUs each. Job completion time was reduced from days to minutes.

⁶Source: Evermann, J. (2016) Scalable Process Discovery using Map-Reduce. *IEEE TSC*, 9 (3), 469-481.
<https://doi.org/10.1109/TSC.2014.2367525>



Source: Evermann, J. (2016) Scalable Process Discovery using Map-Reduce. *IEEE TSC*, 9 (3), 469-481.
<https://doi.org/10.1109/TSC.2014.2367525>

Figure 17.5: Total elapsed time for completion of MapReduce jobs for process discovery

α Algorithm	
Single node	25:00 hours
Medium cluster	1:24 hours
Large cluster	0:08 hours
FHM	
Single node	22:21 hours
Medium cluster	2:01 hours
Large cluster	0:17 hours

Source: Evermann, J. (2016) Scalable Process Discovery using Map-Reduce. *IEEE TSC*, 9 (3), 469-481.
<https://doi.org/10.1109/TSC.2014.2367525>

Table 17.3: Total elapsed time for completion of MapReduce jobs for process discovery

Apache Pig



Apache Pig⁷ is a high-level platform for creating programs that run on Apache Hadoop. It is designed to simplify the complexities of writing low-level MapReduce programs, providing a simpler scripting language called Pig Latin, which abstracts the details of programming from the underlying MapReduce framework. Apache Pig was originally developed at Yahoo to allow analysts using Hadoop to focus more on analyzing large data sets and spend less time having to write Map and Reduce functions. Although it operates at a higher level of abstraction than MapReduce, under the hood, Pig converts these scripts into MapReduce jobs.

Pig Latin is designed to handle all kinds of data, particularly semi-structured data or

⁷<https://pig.apache.org/docs/latest/basic.html>

LOAD	STORE	DUMP	FILTER	DISTINCT
FOREACH... GENERATE	UNION	SAMPLE	JOIN	GROUP
CROSS	ORDER	LIMIT	SPLIT	

Table 17.4: Core Pig Latin operations

datasets that do not conform to a fixed schema. Pig Latin supports operations like join, sort, filter, and combine but is tailored to handle complex nested data structures typical in big data applications. Users can extend the language with their own functions written in Java, JavaScript, Python, etc., to handle custom processing. Typically, Pig Latin programs are about 5-20 times shorter than equivalent Java MapReduce programs. However, Pig Latin remains a procedural language, focusing on how to retrieve or process data, rather than a declarative language like SQL that focuses on what data to retrieve. Table 17.4 shows the core relational operations available in Pig Latin.

To give an impression of Pig Latin, consider the following script⁸ that performs the same word count as the MapReduce example above.

```
input_lines = LOAD 'hamlet.txt' AS (line:chararray);
-- Extract words from each line and put them into
-- a pig bag datatype, then flatten the bag to get
-- one word on each row
words = FOREACH input_lines \
    GENERATE FLATTEN(TOKENIZE(line)) AS word;
-- create a group for each word
word_groups = GROUP words BY word;
-- count the entries in each group
word_count = FOREACH word_groups \
    GENERATE COUNT(words) AS count, group AS word;
-- order the records by count
ordered_word_count = ORDER word_count BY count DESC;
STORE ordered_word_count INTO 'hamlet.out';
```

Apache Hive



https://commons.wikimedia.org/wiki/File:Apache_Hive_logo.svg

Apache Hive is a data warehousing software and SQL-like query language for data stored on Hadoop's HDFS. Hive is designed to make Hadoop accessible to business analysts familiar with SQL, the standard language for relational databases. However, Hive is not a full database. The main focus is to provide data summarization, query, and analysis. While it has similar properties to a database, such as indexing, transactions, and queries, it does not offer real-time queries and row-level updates.

⁸Source: https://en.wikipedia.org/wiki/Apache_Pig

HiveQL⁹ is the SQL-like language that Hive uses. HiveQL includes extensions that allow traditional MapReduce programmers to plug in their custom Map and Reduce functions when it is inconvenient or inefficient to express this logic in HiveQL. Hive reduces the complexity of writing MapReduce jobs by automatically translating HiveSQL queries into MapReduce jobs, allowing users to focus on query statements rather than the complexities of the underlying execution engines.

Consider the following example HiveQL query¹⁰. It implements the same word count as the above Pig Latin script and the earlier MapReduce example. Note the similarities to SQL and the declarative nature. The HiveQL query does not specify what actions to perform, but simply specifies what result to retrieve or select from the data set.

```
DROP TABLE IF EXISTS docs;
CREATE TABLE docs (line STRING);
LOAD DATA INPATH 'hamlet.txt'
OVERWRITE INTO TABLE docs;

CREATE TABLE word_counts AS
SELECT word, count(1) AS count FROM
(SELECT explode(split(line, '\s')) AS word FROM docs) temp
GROUP BY word
ORDER BY word;
```

17.3 Apache Spark



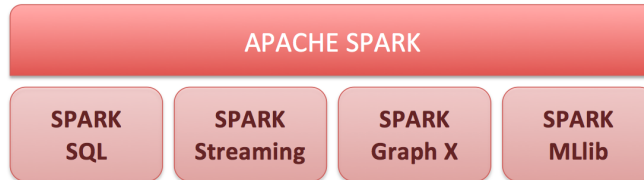
Apache Spark is an open-source, unified analytics system for large-scale data processing. Originally designed at developed at the University of California at Berkeley in 2009, it was donated as an open-source project to the Apache Foundation in 2013. It provides high-level programming interfaces for Java, Scala, Python, and R. It is known for its speed and ease of use in complex analytics across big data. Because of its advantages over Hadoop and MapReduce, it was quickly adopted.

Spark's core feature is its in-memory cluster computing that increases the processing speed of an application significantly over MapReduce, which is disk limited. Spark stores data in RAM across the cluster, which allows it to access this data quickly and speed up the computation times, especially for iterative algorithms common in machine learning and data mining. In contrast to MapReduce, Spark also supports cyclic data flow.

Spark can run on top of existing Hadoop clusters to leverage Hadoop's storage systems, like HDFS or HBase. This allows for easy integration and migration of data-processing

⁹<https://cwiki.apache.org/confluence/display/Hive/LanguageManual>

¹⁰Source: https://en.wikipedia.org/wiki/Apache_Hive



https://commons.wikimedia.org/wiki/File:Sch%C3%A9ma_d%C3%A9tail_outils_spark.png

Figure 17.6: Apache Spark components

tasks between Hadoop components and Spark. Like Hadoop, Spark is designed to scale up from a single server to thousands of machines, each offering local computation and storage. However, Spark can also use cloud-based file storage such as Amazon Web Services S3 or Microsoft Azure storage solutions, and cluster management software other than YARN, such as Mesos or Kubernetes.

Spark provides a unified processing engine that can handle different types of workloads within the same application that have traditionally required separate distributed systems, including batch processing, SQL queries, real-time stream processing, machine learning, and graph processing. This unification reduces management overhead and streamlines data processing pipelines.

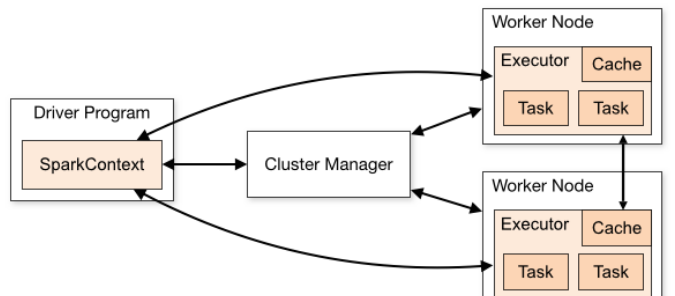
Spark uses a data storage model based on RDDs (Resilient Distributed Datasets), which are automatically rebuilt on cluster node failure. This design ensures that Spark applications can handle node failures on large clusters with minimal impact on data processing tasks.

Spark's easy-to-use programming interface simplify the development of complex, multi-stage data pipelines and sophisticated analytics and machine learning compared to MapReduce. Spark also includes Spark SQL, making it easy to transition from traditional SQL databases to big data processing.

Figure 17.6 shows the components of Spark. Despite its name, Spark SQL also provides more traditional data frame operations, similar to those one might see in the Python Pandas or R Tidyverse packages.

The Apache Spark web site provides a wealth of information, both at the introductory, conceptual level and the advanced, detailed level. The following sections are useful for learning more about Apache Spark:

- [Quick Start](#)
- [SQL, DataFrames and Datasets](#)
- [Structured Streaming](#)
- [Machine Learning](#)



<https://spark.apache.org/docs/latest/img/cluster-overview.png>

Figure 17.7: Apache Spark cluster architecture

Cluster Management

Apache Spark provides its own cluster management, shown in Figure 17.7. An analytics application (“driver program”) uses a Spark context to communicate with the Spark cluster manager. The cluster manager in turn coordinates and controls the worker nodes. Worker nodes run on each computer in the cluster to manage the local resources and execute tasks. When a client application requests execution of a Spark job via the Spark context, the cluster manager assigns worker nodes and their executors to the different tasks in the job. The executors execute the assigned tasks and communicate with the Spark context of the client application for status updates and results.

17.3.1 Spark SQL

Despite the name “Spark SQL”, this component of Spark also offers non-SQL based data management. This section first describes three different kinds of data storage in Spark, introduces execution of transformations and actions on them, then provides some basic examples, and finally illustrates the use of SQL in Spark.

Data Storage Architecture

The fundamental data structure of Apache Spark is the Resilient Distributed Dataset (RDD). RDDs are designed to provide a fault-tolerant, immutable collection of objects that can be processed in parallel. An RDD can contain any type of Python, Java, or Scala data objects. RDDs are immutable, that is, once created they cannot be changed. Data in RDDs is split into logical partitions, which can be processed on different nodes of the cluster. RDDs achieve resilience, that is, fault tolerance, through a lineage graph of the input dataset — each RDD “remembers” how to reconstruct its segments from other datasets by logging the transformations used to build it from other RDDs.

When Apache Spark is run on a Hadoop cluster, RDDs are created from data stored in HDFS. Spark builds the RDD partitions directly from the blocks of the corresponding file stored on the HDFS. This direct alignment allows Spark to leverage HDFS’s loca-

tion model, which places computations near the data. Since RDD partitions correspond to HDFS blocks, operations on RDDs can be scaled across many nodes, allowing Spark to leverage the distributed nature of the HDFS architecture to process large datasets in parallel. The correspondence of Spark RDD partitions and HDFS blocks also means there are two layers of fault tolerance. First, the HDFS replication ensures that data can be recovered in case of node failure from another replica. Second, the Spark RDD lineage information means the RDD can be re-computed from its inputs given its history. RDDs offer a low-level programming interface that is focused on MapReduce. It is procedural and offers no optimization of query operations.

Building on RDDs, Spark provides DataFrames and Datasets that provide a high-level, more abstract programming interface similar to Python Pandas data frames or R dataframes. A DataFrame in Spark is a distributed collection of data organized into named columns. Spark DataFrames use a query optimizer to optimize the execution of DataFrame queries for better performance. DataFrames support various data formats and data sources, including JSON, CSV, and relational databases. DataFrames are also integrated with Spark SQL for running SQL queries.

Datasets in Spark are an extension of DataFrames that provide a type-safe, object-oriented programming interface. Datasets are only available in the Scala and Java programming languages; the Python and R programming languages do not have the type information required to use them.

Spark Execution Principles

Apache Spark's core programming model for DataFrames and Datasets revolves around transformations and actions. These operations adhere to the principle of lazy execution, which is fundamental to Spark's high performance and efficiency in processing large datasets.

Transformations are operations that create a new DataFrame from an existing one. They are considered "lazy", meaning that they do not compute their results right away. Instead, Spark maintains a plan (the "lineage") of all transformations applied to the DataFrame. Common transformations on DataFrames are listed in Table 17.5. Transformations are only executed when an *action* is applied to the DataFrame. This approach allows Spark to optimize data processing, for example by internally rearranging or combining operations.

Actions are operations that trigger computation and return results from a DataFrame to the client application or write results to storage. Examples of actions are shown in Table 17.6. When an action is called on a DataFrame, Spark evaluates the DataFrame transformations that have been built up in the lineage. It then optimizes the execution plan for these transformations and executes the different tasks across the cluster to compute the final results.

<code>select()</code>	Projects a set of expressions and returns a new DataFrame.
<code>filter() or where()</code>	Filters rows using the given condition and returns a new DataFrame with only the satisfying rows.
<code>groupBy()</code>	Groups the DataFrame using the specified columns, so that further aggregation can be performed.
<code>join()</code>	Joins with another DataFrame, using the given join expression.
<code>orderBy() or sort()</code>	Returns a new DataFrame sorted by the specified column(s).
<code>drop()</code>	Removes a column or columns from a DataFrame.
<code>withColumn()</code>	Returns a new DataFrame by adding a column or replacing an existing column that has the same name.
<code>withColumnRenamed()</code>	Renames a column in the DataFrame.
<code>distinct()</code>	Returns a new DataFrame containing the distinct rows of the source DataFrame.
<code>union()</code>	Combines two DataFrames that have the same schema, appending the rows of one DataFrame to another.
<code>cov()</code>	Covariance of columns in a DataFrame.

Table 17.5: Common Spark DataFrame transformations

<code>show()</code>	Displays the contents of the DataFrame in tabular form.
<code>count()</code>	Returns the number of rows in the DataFrame.
<code>collect()</code>	Retrieves the entire DataFrame and returns it as a collection of rows on the driver program.
<code>save()</code>	Saves the DataFrame to an external storage system, such as HDFS or a local filesystem.
<code>take(n)</code>	Returns an array with the first n rows in the DataFrame.
<code>head()</code>	Retrieves the first n rows of a DataFrame.
<code>tail()</code>	Retrieves the last n rows of a DataFrame.
<code>toPandas()</code>	Converts a Spark DataFrame to a Pandas DataFrame, bringing the entire data set into memory.
<code>write.csv()</code>	Writes the contents of a DataFrame to a CSV file.

Table 17.6: Common Spark DataFrame actions

Basic Examples

The examples in this section are intended to illustrate the basic usage of Spark on a Hadoop cluster. They assume that Spark is running on a Hadoop cluster with HDFS

storage. They use Python and the PySpark interactive console.

First, start the local Hadoop cluster (if not already running) and the PySpark console. PySpark is Python with a built-in Spark context object to connect to the cluster manager, as shown in the output below. The Hadoop YARN cluster manager manages the PySpark application, and has a web interface at `localhost:8088`. From there, information about the PySparkShell application can be viewed.

```
sudo systemctl start hadoop.service
pyspark --master yarn
```

The result is the PySparkShell, which is a Python shell with a Spark context object that provides the connection to the cluster manager:

```
Python 3.10.12 (main, Nov 20 2023, 15:14:05) [GCC 11.4.0] on linux
Type "help", "copyright", "credits" or "license" for more information.
Welcome to

   / _ \_ / \_ \_ \_ \_ / \_ \
  / \ \ / - \ / - ^ / \ / ' / \
 / \ / . \ / \ , / \ / / \ / \ \ \
 / \ / \ / \ / \ / \ / \ / \ \ \
                                     version 3.5.1

Using Python version 3.10.12 (main, Nov 20 2023 15:14:05)
Spark context Web UI available at http://10.0.2.15:4040
Spark context available as 'sc' (master = yarn, app id = application_
SparkSession available as 'spark'.
```

The following Python code block reads a file from HDFS and gathers some statistics. Note that the `textFile` is treated as a DataFrame with typical data frame operations such as `count()` and `filter()`. PySpark will provide some information about the job progress on the cluster as these operations are executed.

```
textFile = spark.read.text(
    'hdfs://localhost:9000/user/busi4720/hamlet.txt')
# Number of lines
textFile.count()
# First row
textFile.first()
# How many lines contain the word Hamlet?
textFile.filter(textFile.value.contains("Hamlet")).count()
```

The Python code block below shows another example of Spark DataFrame operations. It finds the longest line by number of words. It imports a number of useful functions from the Spark SQL package. The `sf.split()` function splits a character string on a regular expression into a list of words, `sf.size()` returns the length of that list, `sf.col()` returns a specified columns, and the `sf.max()` function returns the

maximum of its arguments. The `agg()` function specifies an aggregation and the final `collect()` is an action that triggers execution of the transformation operations.

```
# Import useful functions from Spark SQL:
from pyspark.sql import functions as sf
# Split each line and count words as 'numWord',
# then aggregate the 'numWords' columns using 'max':
textFile.select(sf.size(sf.split(textFile.value, "\s+")) \
    .alias("numWords")) \
    .agg(sf.max(sf.col("numWords")))) \
    .collect()
```

The following Python code block implements the word count. It uses two functions from the Spark SQL library, `sf.split()` and `sf.explode()`. The first splits a line of text (a character string) on a regular expression, here, one or more whitespace characters. The `sf.explode()` function converts the list resulting from `sf.split()` into a DataFrame column. This column is then called "word". The `groupBy()` and `count()` operations then aggregate the information. The final `collect()` is an action that triggers actual execution of the transformation operations.

```
# Import useful functions from Spark SQL:
from pyspark.sql import functions as sf

wordCounts = textFile \
    .select(sf.explode(sf.split(textFile.value, "\s+")) \
        .alias("word")) \
    .groupBy("word") \
    .count() \
    .orderBy("count")
wordCounts.collect()
```

Spark Schemas

In Apache Spark, a schema is a structured definition of the columns and their data types in a DataFrame. Schemas serve several important purposes when creating or reading a DataFrame. First, schemas are used to validate the data, ensuring that the data matches the expected format and types. Second, knowing the schema avoids the overhead of inferring data types and enables better optimization techniques.

Spark supports a variety of data types similar to traditional database types, which are used to define the columns in a DataFrame. Common data types in Spark are 'Integer', 'Long', 'Double', 'Float', 'String', 'Boolean', 'Timestamp', 'Date', 'Smallint', 'Tinyint', 'Bigint', and the complex types 'Struct', 'Array' and 'Map'.

A schema in Spark is defined using the Spark schema DDL (data definition language) which looks superficially like an SQL table definition, as shown in the following PySpark example that defines a schema for reading an event log file for process analytics:

```
# Define a schema using Spark schema DDL
logSchema = \
    'caseID STRING, \
     activity STRING, \
     ts TIMESTAMP'
```

The schema can then be used the reading data into a DataFrame, as shown in the Python code block below. A CSV file is read from HDFS using specific options for delimiter and header row and the schema defined above:

```
fname='hdfs://localhost:9000/user/busi4720/eventlog.short.log'

data = spark.read \
    .format('csv') \
    .option('delimiter', '\t') \
    .option('header', 'false') \
    .schema(logSchema) \
    .load(fname)
```

The following PySpark commands provide basic information for the DataFrame:

```
data.printSchema()
data.count()
data.show(5)
data.summary().show()
```

Spark SQL

The word count example above used DataFrame operations similar to those in R or Pandas on Spark DataFrames. Another way to work with DataFrames is to treat them as a table and use SQL queries.

DataFrames can be turned into temporary tables, called "views". These are not physically recorded and are lost when the PySpark application ends (or the view is explicitly destroyed).

```
data.createOrReplaceTempView('log')
```

Alternatively, a DataFrame can written to a permanent table. This table persists in Hadoop storage beyond the PySpark application:

```
data.write.saveAsTable('log_table')
```

In either case, the `spark.sql()` function allows the use of SQL commands on the temporary or permanent table:

```
result_df = spark.sql('select * from log limit 5')
result_df.show()
```

For a more elaborate example, consider the construction of a Directly-Follows-Graph (DFG), as used in business process analytics. This is relatively easy to do with the following SQL query. It computes the number of times an activity directly follows another activity, as well as the mean time between two activities following each other.

```
sql_query = \
    'SELECT COUNT(*), l1.activity AS activity1, \
     l2.activity AS activity2, AVG(l2.ts - l1.ts) AS dtime \
      FROM log AS l1 JOIN log AS l2 ON l1.caseid=l2.caseid \
     WHERE l2.ts = (SELECT MIN(ts) FROM log l3 \
      WHERE l3.caseid=l1.caseid AND l3.ts > l1.ts) \
      GROUP BY GROUPING SETS((l1.activity, l2.activity))'
```

Query execution may require some time due to the multiple self-joins on the log table.

```
# Run the query, show the results
dfg = spark.sql(sql_query)
dfg.count()
dfg.show()
```

To illustrate the concept of a query plan and query plan optimization, it may be useful to examine the optimized query plan that Spark uses for the actual calculations. The output provides information about the logical query plans, both the initial and the optimized one, as well as the physical query plan, the set of tasks that Spark runs on the cluster nodes to calculate the result.

```
# Explain the query plan:
dfg.explain(mode='formatted')
dfg.explain(True)
```

So far, the Spark examples have been run interactively from the PySpark console. However, longer running analytics jobs are typically submitted to the Spark cluster for execution as self-contained applications so the analyst does not have to wait for results to be returned. To do this, one can use the `spark-submit` command and specify the PySpark file and its arguments. The following example downloads a Python application file (which contains the above code to compute the DFG) and then submits it to the cluster manager for execution, providing the HDFS file name as its argument.

Result will be written to HDFS. While the job is running, use Hadoop Job Tracker at <https://localhost:8088> to track the status of nodes and the progress of jobs.

```
# Download file
wget https://evermann.ca/busi4720/spark_dfg.py
# Submit to Spark/Hadoop cluster
spark-submit --master yarn spark_dfg.py \
  hdfs://localhost:9000/user/busi4720/eventlog.short.log
```

17.3.2 Spark Machine Learning

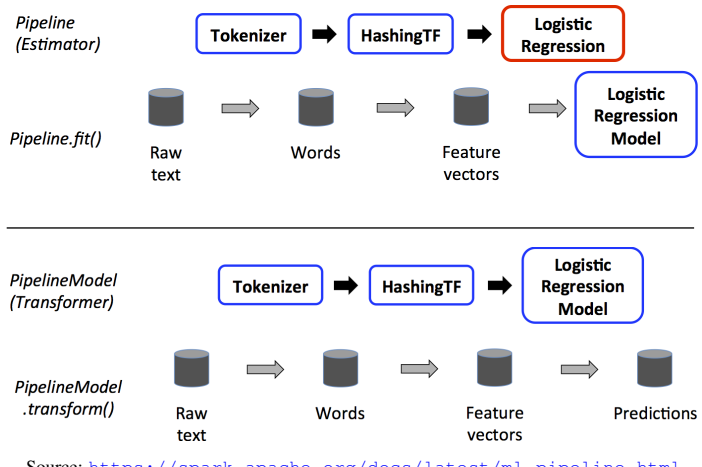
Apache Spark provides support for machine learning through its MLlib and Spark ML libraries. Both libraries offer common learning algorithms like classification (logistic regression, decision trees, support vector machines, random forests, etc.), regression (generalized linear regression, regression trees, etc.), and clustering (for example, k-means). Utilities for feature extraction, transformation, dimensionality reduction (for example, PCA), and feature selection are provided to prepare data for machine learning models. MLlib and Spark ML support the concept of pipelines, which simplify the process of transforming data and tuning parameters. Models and algorithms can be saved to and loaded from storage, facilitating the reuse and application of models across different applications and frameworks. Spark ML provides a higher-level programming interface built on DataFrames for constructing ML pipelines, whereas MLlib is the original machine learning library for Spark and is based on RDDs. Spark ML is the primary focus of development; while still supported, MLlib receives less attention in terms of new features or performance improvements.

In Apache Spark ML, the concepts of pipelines, transformers, and estimators are central to defining data transformations and machine learning workflows in a reusable, manageable way. A *pipeline* in Spark ML represents a workflow that consists of a sequence of stages, each of which is either a *transformer* or an *estimator*. These stages are executed in order and transform the input DataFrame as it passes through each stage. Once a pipeline is defined, it acts like an estimator. Running the pipeline's `fit()` method on a DataFrame produces a fitted model ("PipelineModel"), which is a Transformer.

Transformers take a DataFrame as input and return a new DataFrame with more features or a different arrangement of features as output. Common examples of transformers are the 'Tokenizer' that splits text into words, the 'StringIndexer' that converts a column of labels to a column of integer indices, or the 'VectorAssembler' that combines multiple columns into a single vector.

An *Estimators* represents a learning algorithms that fits or trains on data. Estimators provide a `fit()` method, which accepts a DataFrame and produces a model, which in turn is a Transformer. Common examples of Estimators include 'LogisticRegression', 'DecisionTreeClassifier' or "KMeans".

Figure 17.8 illustrates the use of Spark ML pipelines. The upper panel shows a pipeline consisting of two transformers (blue), followed by an estimator (red). Once the `fit()`



Source: <https://spark.apache.org/docs/latest/ml-pipeline.html>

Figure 17.8: Pipelines in Spark ML

method of the pipeline is called, the estimator (and the entire pipeline) become a transformer (blue). The lower panel shows that this fitted/trained transformer pipeline can then be used to transform input into predictions.

The following example illustrates the use of Spark ML for binary classification. The example requires some feature transformation to illustrate the use of the pipelines. The full example is available to run as self-contained application¹¹ using spark-submit on a Spark cluster. Remember that the Hadoop job tracker¹² is available to track the status of the submitted Spark job. The data set for this example is a public dataset originally from the UCI machine learning laboratory¹³.

```
# Download file
wget https://evermann.ca/busi4720/spark_ml.py

# Get the dataset and put on HDFS
wget https://evermann.ca/busi4720/mushrooms.csv
hdfs dfs -put mushrooms.csv

# Submit to Spark/Hadoop cluster
spark-submit --master yarn spark_ml.py \
    hdfs://localhost:9000/user/busi4720/mushrooms.csv
```

The remainder of this section illustrates the code in detail. First, the schema is defined for reading the CSV data set into a Spark DataFrame and to validate the data types:

¹¹https://evermann.ca/busi4720/spark_ml.py

¹²<https://localhost:8080>

¹³Source: <https://archive.ics.uci.edu/dataset/848/secondary+mushroom+dataset>, CC-BY 4.0 license

```
the_schema = 'class STRING, `cap-diameter` DOUBLE, \
`cap-shape` STRING, `cap-surface` STRING, \
`cap-color` STRING, `does-bruise-or-bleed` STRING, \
`gill-attachment` STRING, `gill-spacing` STRING, \
`gill-color` STRING, `stem-height` DOUBLE, \
`stem-width` DOUBLE, `stem-root` STRING, \
`stem-surface` STRING, `stem-color` STRING, \
`veil-type` STRING, `veil-color` STRING, \
`has-ring` STRING, `ring-type` STRING, \
`sparse-print-color` STRING, habitat STRING, \
season STRING'
```

The `spark.read()` function uses the schema and specific parameters for delimiter and header rows to read the CSV file into a Spark DataFrame. The "veil-type" column is then dropped and missing values are replaced by the character string 'NULL'.

```
fname='hdfs://localhost:9000/user/busi4720/mushrooms.csv'

data = spark.read \
    .format('csv') \
    .option('delimiter', ',') \
    .option('header', 'true') \
    .schema(the_schema) \
    .load(fname)
data = data.drop('veil-type')
data = data.fillna('NULL')
```

Next, import the required packages:

```
# Import all required pieces:
from pyspark.ml import Pipeline
from pyspark.ml.classification import LogisticRegression
from pyspark.ml.feature import StandardScaler, \
    StringIndexer, OneHotEncoder, VectorAssembler
from pyspark.ml.evaluation import BinaryClassificationEvaluator
from pyspark.ml import PipelineModel
```

The numerical feature columns are concatenated ("assembled") into a single vector column using the `VectorAssembler` transformer:

```
numFeatures = VectorAssembler(
    inputCols = ['cap-diameter', 'stem-width', 'stem-height'],
    outputCol = 'numFeatures')
```

The `StandardScaler` transformer scales and standardizes the numerical features:

```
scaler = StandardScaler(inputCol='numFeatures',
                        outputCol='numFeaturesS')
```

The categorical variables are one-hot encoded using dummy variables. First, identify the columns with 'string' data type, and construct column names for the indexed and one-hot encoded corresponding columns:

```
categoricalCols = \
    [name for (name, dtype) in data.dtypes if dtype=='string']
indexOutputCols = [x + 'index' for x in categoricalCols]
oheOutputCols = [x + 'ohe' for x in categoricalCols]
```

A `StringIndexer` is a transformer that transforms a column or set of columns of character strings into a column or set of columns of integer indices.

```
stringIndexer = StringIndexer(
    inputCols = categoricalCols,
    outputCols = indexOutputCols,
    handleInvalid='skip')
```

The `OneHotEncoder` takes one or more numerical category index columns and returns the corresponding one-hot encoded columns:

```
oheEncoder = OneHotEncoder(
    inputCols = indexOutputCols,
    outputCols = oheOutputCols)
```

Finally, the numerical and categorical feature columns are combined into a column with the entire feature vector using another `VectorAssembler` transformer:

```
# Assemble all features into a feature vector
vecAssembler = VectorAssembler(
    inputCols = oheOutputCols+['numFeaturesS'],
    outputCol = 'feature_vec')
```

The targets are also encoded as numerical indices, using the `StringIndexer` transformer:

```
# Encode the target classes as numbers
stringIndexTarget = StringIndexer(
    inputCols = ['class'],
    outputCols = ['classIndex'],
    handleInvalid='skip')
```

As the last required element, the `LogisticRegression` estimator is defined, accepting the feature vector column and the target column:

```
# Create the classification estimator
logReg = LogisticRegression(
    featuresCol = 'feature_vec', labelCol = 'classIndex')
```

All components can be assembled into a pipeline:

```
# Put all components into the pipeline
pipeline = Pipeline(stages=[
    numFeatures,
    scaler,
    stringIndexer,
    oheEncoder,
    vecAssembler,
    stringIndexTarget,
    logReg])
```

To train the model, the data set is split into training and test data, using the `randomSplit` transformation on the Spark DataFrame:

```
# Create train/test data split
train_data, test_data = data.randomSplit([.66, .33], seed=1)
```

Calling the pipeline's `fit` method will turn the final estimator, and the entire pipeline into a fitted model, that is, a transformer:

```
# Fit the model to the training data
pipelineModel = pipeline.fit(train_data)
```

Performance metrics for the training data can be retrieved from the last stage of the pipeline, that is, from the `LogisticRegression` estimator (now a transformer):

```
# Summary of the training data performance
summary = pipelineModel.stages[-1].summary
summary.accuracy
summary.areaUnderROC
summary.fMeasureByThreshold.show()
summary.precisionByLabel
summary.recallByLabel
summary.roc.show()
```

Fitted estimators, including whole pipelines, become transformers. The `transform()` method of a transformer can be used to transform inputs into output predictions:

```
trainPred = pipelineModel.transform(train_data)
testPred = pipelineModel.transform(test_data)
```

Apache Spark ML provides "evaluators" to evaluate the predictive performance of models. The `BinaryClassificationEvaluator` focuses on the AUC. It accepts the true classes and can then evaluate the predictions created by the pipeline against the true class indices:

```
# Evaluate the model using AUC
evaluator = BinaryClassificationEvaluator(
    labelCol='classIndex')
evaluator.evaluate(trainPred)
evaluator.evaluate(testPred)
```

To help manage trained models for later reuse, Spark ML provides methods to save and load fitted models/pipelines:

```
# Save the fitted model for later re-use:
pipelineModel.write().overwrite().save('myFirstModel')

# Load a saved model:
savedModel = PipelineModel.load('myFirstModel')
```

17.4 Stream Analytics

So far, this section has focused on what is known as batch processing or batch analytics. This involves collecting data over a period of time, working with a finite, but possibly large data set, and processing it in large batches at a scheduled time, e.g. daily, weekly, or monthly.

Stream analytics on the other hand focuses on data that cannot be, or does not need to be stored permanently, typically due to high data volume and high data velocity. For example, many industrial systems are extensively instrumented with sensors that provide readings every millisecond. Storing terabytes or petabytes worth of low-level detailed data every day is infeasible. Moreover, decision making based on the data may need to happen continuously and in real-time. Stream analytics focuses on processing such data "on-the-fly", as it is flowing through the analytics application, without being able to store it, and without being able to recall it – once it's passed through, it's gone. Data is continuously processed in real-time and there may be perpetual stream of input data that needs to be processed.

Example use cases in business analytics are customer click-stream analysis for real-time pricing on web sites, machine sensor data processing for failure warnings or

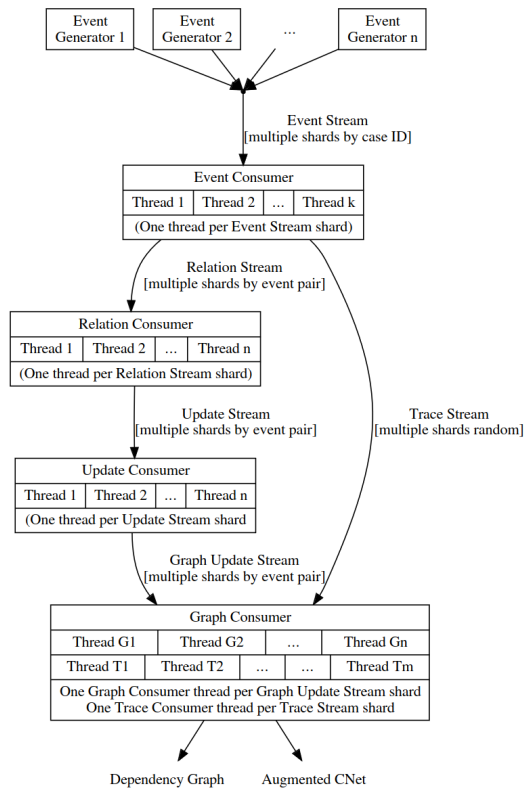
alarms, financial transaction fraud monitoring, financial market data and financial news analysis, or business process compliance monitoring. All these applications have the same characteristics: Analysis must happen in real time, as the data is ingested; the flow of incoming data never stops; and there is too much data to be stored for batch analytics.

Stream analytics applications are often conceptualized as a network of processing nodes, each node ingests a data stream, performs fast, low-latency processing, and emits another data stream that is sent or piped to another set of nodes as input.

To illustrate this, consider the use case of continuous business process discovery for streaming data¹⁴. The flexible heuristics miner is implemented for streaming data as a network of processing nodes. The system ingests activity completion events from process-aware information systems and, as each event is ingested, continuously updates the discovered business process model in real-time. The system was implemented on Amazon Web Services Kinesis. It uses multiple data streams between processing nodes. Each data stream is essentially a queue into which data is put at one end and read in the same order from the other end, but data is never stored. The data processing nodes perform the various computations required to construct the process model and, for They are implemented using multipled threads/executors for performance. Figure 17.9 shows the system architecture.

Running on 5 compute nodes, the system is able to process over 5 million events per minute, or over 150,000 complete process traces per minute and continuously updates the discovered process model. Figure 17.10 shows the data throughput for each of the data streams in the system during a 3-hour period; the vertical axis is in millions of records processed.

¹⁴Source: Evermann, J., Rehse, J.-R., and Fettke, P. (2016) Process Discovery from Event Stream Data in the Cloud - A Scalable, Distributed Implementation of the Flexible Heuristics Miner on the Amazon Kinesis Cloud Infrastructure. *CloudBPM Workshop on Business Process Monitoring and Performance Analysis in the Cloud at the 8th IEEE International Conference on Cloud Computing Technologies and Science (CloudCom 2016)*



Source: Evermann, Rehse, Fettke (2016)

Figure 17.9: Flexible Heuristics Miner implemented on AWS Kinesis, system architecture



Source: Evermann, Rehse, Fettke (2016)

Figure 17.10: Flexible Heuristics Miner implemented on AWS Kinesis, data throughput

17.5 Spark Streaming

Apache Spark Streaming enables scalable, high-throughput, fault-tolerant stream processing of live data streams. It can ingest data from various sources like Kafka, Flume, Kinesis, or network sockets, and process it using algorithms expressed with high-level functions like map, reduce, join, and window. Spark Streaming can provide output to a number of destinations, such as distributed file systems, databases, and visual dashboards (Figure 17.11).

Spark Streaming processes live streams of data in small batches, known as *micro-batches*, containing one or a few data records. This allows the framework to achieve high throughput and low latency. These micro-batches are created at user-defined intervals and processed by the Spark engine to generate the final stream of results in batches (Figure 17.12).

Spark Streaming represents data streams as unbounded tables (Figure 17.13). With this representation Spark provides a unified programming model for both batch and stream processing, using the same Spark DataFrame operations for both modes. Spark Streaming can be combined with other big data tools supported by Spark, such as Spark SQL, MLLib for machine learning, and GraphX for graph processing. This integration allows for seamless data processing pipelines that include streaming data, batch data, and interactive queries.

Spark Streaming supports stateful computations, allowing it to *Maintain state* across different batches of data. This is crucial for applications that require tracking session information or aggregating data over time. Spark Streaming also provides capabilities for *windowed computations*, where data transformations are applied over a sliding window of data. This is essential for tasks that need to group and aggregate data over specific time frames.

Apache Spark Streaming provides various options for controlling the timing of streaming data processing through its trigger modes, and different ways of managing output data via output modes. *Triggers* in Apache Spark Streaming determine how often the streaming data should be processed. There are several types of triggers:



<https://spark.apache.org/docs/latest/img/streaming-arch.png>

Figure 17.11: Spark Streaming input sources and output destinations

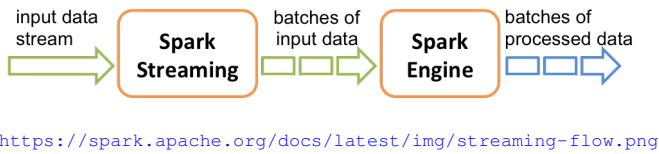


Figure 17.12: Speark Straming batches

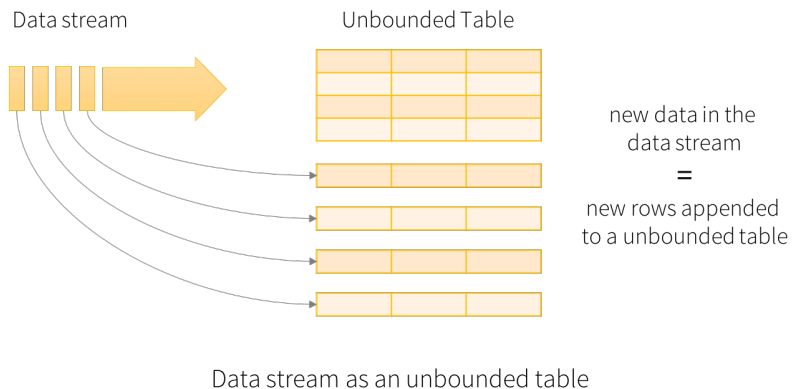


Figure 17.13: Spark data stream as an unbounded table

- *Micro-batch*: The next micro-batch is processed as soon as the previous micro-batch is completed. This mode aims to achieve the highest possible throughput by keeping the system continuously busy.
- *Fixed time interval*: This trigger processes micro-batches at user-specified fixed intervals, such as every 5 seconds. This is useful for scenarios where data should be processed at regular, predictable intervals.
- *Once*: A one-time trigger processes a single batch in response to an event or as a one-off computation. This can be useful for testing or for updating results at irregular intervals.
- *Continuous*: In continuous processing mode, Spark processes records immediately as they arrive, which significantly lowers the end-to-end processing time (latency). This mode is still experimental and may not support all the features of structured streaming.

Output modes in Spark Streaming dictate what gets written to the output destination at the end of each trigger interval. There are several output modes to choose from:

- *Complete Mode*: In this mode, the entire updated result table is outputted after every trigger. It is suitable for scenarios where the full snapshot of the table is needed after each processing interval.
- *Append Mode*: This mode is used when you only want to output the rows that were added to the result table since the last trigger. It is commonly used for cases where only new data is of interest (e.g., new records from a stream).
- *Update Mode*: The update mode outputs only the rows that were updated in the result table since the last trigger. It does not output the rows that have not changed, making it more efficient than complete mode if only changes are necessary.

Spark Streaming is integrated with Spark ML machine learning. It provides streaming implementations of basic machine learning models (linear regression, logistic regression, k-means clustering) that can be trained on continuous data streams. Spark Streaming also offers the ability to predict from data stream for models that were trained off-line, for models created by Spark ML.

Example

To illustrate Spark Streaming, the following PySpark example implements a real-time word count system that ingests data from a network connection ("socket")¹⁵.

The first Python code block imports all necessary functions:

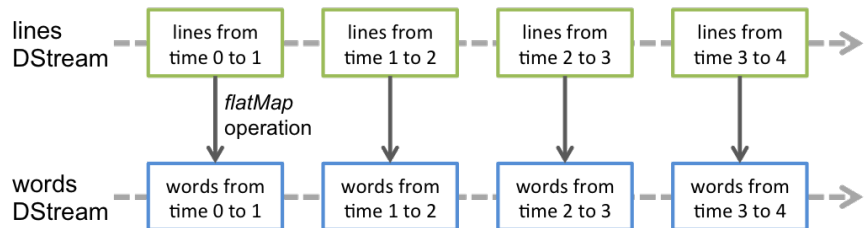
```
from pyspark.sql.functions import explode, split, col, desc, \
    window, current_timestamp
```

The next code block creates a stream reader to read from a network socket on the local machine on port number 9999. The result, `lines`, is a Spark DStream object representing the input data stream. Note that the stream reader opens a *client* socket, i.e. the socket must already have been opened for writing by the data producer, otherwise the `load()` action will fail.

```
lines = spark.readStream \
    .format('socket') \
    .option('host', 'localhost') \
    .option('port', 9999) \
    .load()
```

Next, processing of individual lines is done, using the same functions as in the earlier Spark DataFrame examples. In the following block of Python code `words` is another DStream that is connected to the `lines` DStream, as illustrated in Figure 17.14.

¹⁵Source: <https://spark.apache.org>



<https://spark.apache.org/docs/latest/img/streaming-dstream-ops.png>

Figure 17.14: Spark Streaming line and word DStreams

```
words = lines.select(explode(split(col('value'), '\\s')).alias('word'))
```

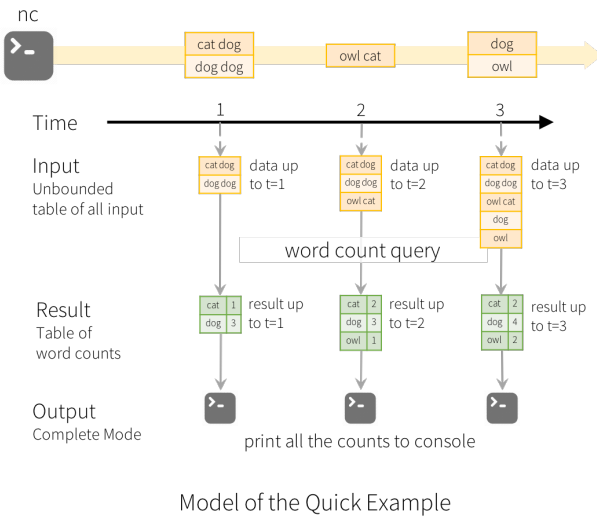
Individual words are then processed, again, in the same way as in the earlier Spark DataFrame example. The following Python code block creates `counts` as another Dstream, connected to the `words` DStream.

```
counts = words.groupBy('word') \
    .count() \
    .sort(desc('count'))
```

Finally, an output writer is defined with a "complete" output mode that writes the complete result after each micro-batch, and a 5 second timer interval trigger on processing – Spark Streaming will read and process a micro-batch every 5 seconds. The checkpoint location is where Spark Streaming stores information about the status of the data streams so it can recover in case processing is interrupted. This ensures fault tolerance, and allows Spark Streaming to guarantee that every record is processed and every record is processed only once. Figure 17.15 illustrates the complete example.

```
writer = counts.writeStream \
    .format('console') \
    .outputMode('complete') \
    .trigger(processingTime='5 second') \
    .option('checkpointLocation', \
        'hdfs://localhost:9000/user/busi4720/')
```

To see this Spark Streaming application in operation, use a Bash command shell to open a network socket using the `nc` command:



Model of the Quick Example

<https://spark.apache.org/docs/latest/img/structured-streaming-example-model.png>

Figure 17.15: Spark Streaming complete word count example

```
nc -kl 9999
```

Then, start the stream data processing by starting the writer. This returns a streaming query object that provides progress information. The "start()" method is a non-blocking operation so that stream processing will occur in the background.

```
streamingQuery = writer.start()
```

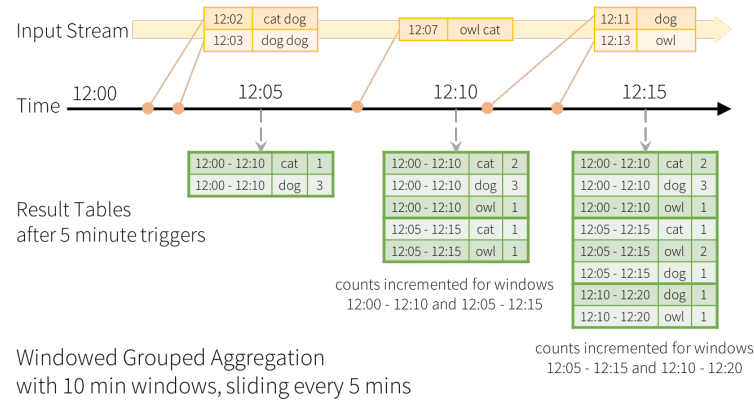
The query object can be used to get progress information, through its `lastProgress` attribute:

```
print(streamingQuery.lastProgress)
```

The query object also provides a `stop()` method to end the processing:

```
streamingQuery.stop()
```

This basic example can be extended to illustrate time windowing, illustrated in the following Python code block. First, the line processing is changed to not only split



<https://spark.apache.org/docs/latest/img/structured-streaming-window.png>

Figure 17.16: Spark Streaming time windowing

lines into words, but also to record the event timestamp. That is, the `words` DStream will contain two columns, `word` and `eventTime`:

```
words = lines \
    .select(explode(split(col('value'), '\\s')).alias('word')) \
    .withColumn('eventTime', current_timestamp())
```

The following Python code block changes the definition of the `counts` DStream to group by words but now for one minute windows of the `eventTime` column; windows are updated every 30 second, that is, overlapping time windows. Figure 17.16 illustrates the time windowing concept.

```
counts = words \
    .groupByKey('word', window('eventTime', '1 minute', '30 second')) \
    .count() \
    .sort(desc('count'))
```

17.6 Review Questions

Introduction

1. Define big data analytics and discuss its significance to business practices.
2. What are the three main "V's of big data? Explain how they characterize big data challenges and opportunities.

3. The text mentions two additional Vs, veracity and value. Describe these concepts and explain their importance in the context of big data.
4. Why is timely data processing important in scenarios involving high-velocity data? Provide an example where delay in data processing could be detrimental.
5. What are the implications of poor data veracity? How can organizations ensure the accuracy and reliability of their data?
6. Discuss the concept of value in big data. What steps must organizations take to transform big data into actionable insights?
7. What are some challenges organizations face when scaling from desktop data analysis tools to industrial-scale big data analytics platforms?

Apache Hadoop

8. Describe the primary function of Apache Hadoop and its origins.
9. Explain the concept of 'data locality' in the context of Hadoop. Why is it beneficial to process data where it is stored?
10. What are the three main components of Hadoop? Briefly describe the role of each component.
11. Discuss the reliability features of Hadoop. How does Hadoop ensure data is not lost in case of a node failure?
12. Describe the process of data replication in HDFS. Why is data replicated across different nodes, and how does this impact system performance?
13. Explain the architecture of HDFS. What are the roles of the NameNode and DataNodes?
14. What are the limitations of the Hadoop ecosystem, and how have new developments or additional tools addressed these challenges?
15. Reflect on how the principles of Hadoop (for example, data locality and redundancy) apply to practical scenarios such as disaster recovery and data processing efficiency.
16. Discuss the impact of Hadoop's 'write once, read many times' model on data analysis tasks. What types of applications are best suited for this model?

MapReduce

17. Define the MapReduce programming model and explain its primary purpose in handling large data sets.
18. Describe the roles of the Map and Reduce functions in a MapReduce job. What types of operations might each perform?
19. Explain the process of data flow from input to output in a MapReduce job, including the stages of Map, Shuffle, and Reduce.
20. Discuss how MapReduce utilizes the distributed storage provided by HDFS for both computation and storage of intermediate results.
21. What are the limitations of the MapReduce model regarding the types of data flows it supports? Why does it struggle with iterative or cyclic data flows?
22. Describe the roles of the Resource Manager and NodeManager in the YARN architecture. How do they interact during the execution of a MapReduce job?

23. What happens during the shuffle phase of a MapReduce job? Explain how data is distributed and prepared for the Reduce phase.
24. Explain the role of the Application Master in a MapReduce job executed on a YARN cluster. How does it manage job execution and resource allocation?
25. How does the number of Reduce instances relate to the number of unique key values in the input data? What determines the number of Reduce tasks in a job?
26. What is the purpose of the Map function in the MapReduce model? Provide an example of a simple Map function.
27. Explain the concept of key-value pairs in the context of MapReduce. How are these used throughout the MapReduce process?
28. Discuss how the Reduce function aggregates the outputs from the Map function. Provide an example where this aggregation is critical to the outcome of the MapReduce job.
29. Why might intermediate data in a MapReduce job be larger than the input data? What challenges does this present?

Apache Pig

30. What is Apache Pig?
31. Explain the main features of Pig Latin and how it simplifies writing data analysis programs compared to MapReduce.
32. What are some of the core operations in Pig Latin? Provide examples of how two of these operations can be used in data processing.
33. Compare and contrast the procedural nature of Pig Latin with the declarative nature of SQL. What are the implications of each approach for data analysis?

Apache Hive

34. What is Apache Hive and what need does it fill within the Hadoop ecosystem?
35. What are the benefits of Hive transforming HiveQL queries into MapReduce jobs?
36. How do Hive and Pig differ in terms of their approach to handling data on Hadoop? Consider aspects such as ease of use, flexibility, and the type of abstraction they provide.

Apache Spark

37. What is Apache Spark and why is it considered a unified analytics system?
38. What is the primary reason for Apache Spark's fast adoption in the industry compared to Hadoop's MapReduce?
39. Explain the concept of in-memory cluster computing in Spark. How does this improve processing speed compared to disk-based systems like MapReduce?
40. How does Spark integrate with existing Hadoop clusters and why is this beneficial for users already using Hadoop?
41. Describe the various types of workloads that Spark's unified engine can handle. How does this versatility affect system management and efficiency?

42. What are Resilient Distributed Datasets (RDDs)? Discuss their importance in Spark's architecture, including how they handle failures.
43. Describe the concept of data lineage in Spark's architecture. How does it assist in the re-computation of RDDs if part of the data or process is lost?
44. Explain the role of Spark SQL within the Apache Spark ecosystem. How does it integrate traditional SQL database functionality with big data processing?
45. Discuss the execution principles of Apache Spark, focusing on transformations and actions. What does lazy execution mean and why is it beneficial?
46. How does Apache Spark manage large-scale data processing across a cluster? Explain the roles of the driver program, cluster manager, and worker nodes.
47. What is a schema in the context of Apache Spark, and what purposes does it serve?
48. Describe a scenario where using SQL to operate on a DataFrame could be more advantageous than using DataFrame API methods directly.

Apache Spark Machine Learning

49. Explain the concept of a machine learning pipeline in Spark ML. How does it enhance the workflow of machine learning models?
50. What is a *Transformer* in Spark ML, and what role does it play in a machine learning pipeline? Provide examples.
51. Describe what an *Estimator* is and its function within the Spark ML framework. Include examples of common estimators.
52. How does Spark ML leverage the concept of lazy execution in its machine learning pipelines?
53. Detail how a fitted machine learning pipeline acts as a transformer. What does this imply for new input data?

Apache Spark Streaming

54. Define stream analytics and contrast it with batch processing. What are the key characteristics that differentiate the two?
55. Explain why stream analytics is essential for data with high volume and velocity. Provide examples of scenarios where stream analytics is preferable.
56. Explain the importance of real-time decision-making in stream analytics. How does this impact the design of stream processing systems?
57. Describe the unified programming model of Spark Streaming. How does it enable seamless transition between batch and stream processing?
58. Explain the concept of micro-batching in Spark Streaming. How does it contribute to achieving high throughput and low latency?
59. Discuss the different trigger modes available in Spark Streaming. What are the use cases for each mode?
60. What are the output modes available in Spark Streaming and how do they differ? Provide scenarios in which each would be used.
61. Analyze the benefits and limitations of using continuous processing mode in Spark Streaming. What types of applications benefit most from this mode?

Chapter 18

Reinforcement Learning – Tabular Methods

Sources and Further Reading

The material in this chapter is based on the following sources.

Richard S. Sutton and Andrew G. Barto (2018) *Reinforcement Learning – An Introduction*. 2nd edition, The MIT Press, Cambridge, MA. (SB)
<http://incompleteideas.net/book/the-book.html>

Chapters 2–7

(CC BY-NC-ND License)

The Sutton & Barto book is a standard introductory textbook on reinforcement learning and widely used. It is very approachable, but at the same time also detailed and thorough in its exposition. Its focus is on RL prior to the use of neural networks for function approximation, so up to about 2015. While it does not provide Python code itself, the pseudo-code in the book is easily implemented.

Resources

Complete implementations of all examples in this chapter are available in the following GitHub repo:

<https://github.com/jevermann/busi4720-rl>

The project can be cloned from this URL:

<https://github.com/jevermann/busi4720-rl.git>

18.1 Introduction

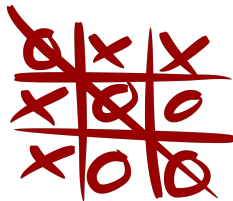
Reinforcement learning (RL) is a type of machine learning in which learning *agents* operate in an *environment* by taking *actions* and receiving *rewards*. The aim is to learn *optimal policies*, that is, those actions for each state that will *maximize* the sum of future rewards. The agent discovers which actions to take and how useful or valuable they are in each state by trying them and observing the reward and the new state.

Initially, agents have little or no knowledge of their environment, so most of the actions will be random exploratory actions. As agents learn more about their environment, they will want to exploit this knowledge by taking the valuable actions, rather than exploring randomly. On the other hand, less exploration may also mean that better actions will not be discovered. In other words, an agent is faced with a trade-off between exploration and exploitation.

What makes RL challenging is both the incomplete knowledge of the environment as well as the stochastic nature of the environment. Taking the same action in the same state will not always yield the same reward, and will not always put the agent in the same new state. The lack of complete knowledge of the environment also means that typical RL problems cannot be solved by optimization; optimization requires full knowledge of the environment, which for stochastic environments, includes knowledge of any probability distributions. This requirement is not fulfilled in RL problem settings.

The core elements of an RL problem are the following:

- **Policy π :** A deterministic policy π specifies for each state s the action to take, whereas a stochastic policy π specifies for each state s a probability distribution over the possible actions a in state s .
- **Reward R :** The reward is received from the environment after each action. The reward may be positive, negative, or zero. In designing RL problems, the reward function is critical to inducing the correct learning behaviour and having the RL agent solve the right problem.
- **Return G :** The return is the possibly discounted sum of future rewards. The discount factor $0 < \gamma \leq 1$ expresses the fact that immediate rewards are worth more than future rewards. This is due to the uncertain nature of future rewards. In a stochastic environment of which the agent has incomplete knowledge, future rewards may or may not accrue as expected.
- **State value function v :** This function expresses how valuable it is for an agent to be in any particular state s . It is defined as the expected return for each state.
- **Action value function q :** This function expresses how valuable it is for an agent in a particular state s to take a specific action a . It is defined as the expected return for each state and action taken in that state.
- **Model p :** This is a set of probability distributions over rewards and new states for every pair of current state s and action a . It expresses the stochastic behaviour of the environment. If an RL agent had such a model of the environment, an optimal



https://commons.wikimedia.org/wiki/File:Tic_tac_toe.svg

Figure 18.1: The game of Tic-Tac-Toe

policy can be found using dynamic programming, a type of optimization. RL agents typically do not try to build such a model, but instead focus on learning the state value function, the action value function, or the policy directly.

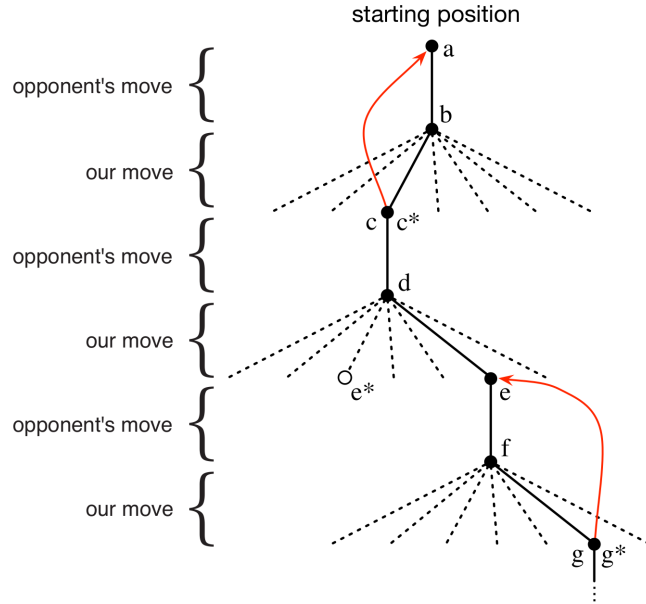
Introductory Example

Consider an RL agent learning the game of Tic-Tac-Toe ("naughts-and-crosses"), as shown in Figure 18.1. A state is defined as the position of all the X and O on the board; the possible actions in each state are to place an X in a free space (assume the agent plays X). The reward at each step is 0 except it is +1 when the game is won. Clearly, the value of any state with a row of X-X-X is 1 because the reward in this case is 1. The value of any state with a row of O-O-O or a full board is 0 because no future reward can occur.

Figure 18.2 indicates a sequence of moves by the RL agent and the opponent where a starred state (e.g. c^*) indicates an optimal state. Beginning from state a , the opponent makes the first move and brings the agent to state b . The agent then exploits the knowledge about the environment that is reflected in its policy or state value function and chooses the optimal action to move to state c^* . The opponent's move leads to state d . Now the agent makes an exploratory move and rather than moving to optimal state e^* it moves to state e . The opponent moves the state to f and the following action of the agent is exploiting behaviour again.

Behaviour that exploits knowledge about the environment, that is, the current action value function and policy, is called *greedy* behaviour as it seeks to maximize the value of the next state. In contrast, exploratory behaviour is typically *random* behaviour.

After each greedy action from state s_t , the RL agent updates its value function for the state s_t based on the value of the new state s_{t+1} . The intuition is that if the optimal action from an initial state of low value results in a new state of high value then the value of the initial state should reflect this. In other words, the updated value of the initial state should be closer to that of the final state. This leads to the central *update rule* in *temporal-difference learning*:



Source: SB Figure 1.1

Figure 18.2: Exploration and exploitation in an RL environment

$$V(S_t) \leftarrow V(S_t) + \alpha [V(S_{t+1}) - V(S_t)] \quad (18.1)$$

Here, V is the value function, and α is a *step size* that determines the rate of update or learning. The term $V(S_{t+1}) - V(S_t)$ is called the *error* and the term $V(S_{t+1})$ is called the *update target*.

Applications in Business and Management

There are many applications for reinforcement learning in business and management. Consider the following examples:

- *Dynamic Pricing*: Dynamic pricing involves setting flexible prices for products or services based on current market demands. RL algorithms can help businesses optimize pricing strategies in real-time by learning from consumer behavior and competitor actions, maximizing revenue or market share.
- *Supply Chain Optimization*: In supply chain management, RL can optimize inventory levels, improve logistics, and manage the supply chain network's dynamic environment. By learning from historical data and ongoing operations, RL algorithms can make adjustments to inventory and shipping strategies, reducing costs and improving service levels.



https://commons.wikimedia.org/wiki/File:Antique_one-armed_bandit,_Ventnor,_Isle_of_Wight,_UK.jpg

Figure 18.3: An "one-armed bandit" slot machine

- *Customer Interaction Management:* Reinforcement learning can enhance customer relationship management systems by learning to tailor interactions based on customer behavior. This includes optimizing marketing strategies, personalizing recommendations, and improving customer service, all aimed at enhancing customer satisfaction and loyalty.
- *Financial Portfolio Management:* In finance, RL can be used for portfolio management, where the goal is to optimize the allocation of assets in a portfolio over time. RL algorithms can adapt to changes in market conditions, learning to maximize returns or minimize risk based on the investment strategy.
- *Manufacturing Process Optimization:* RL algorithms can be applied to control and optimize manufacturing processes by continuously learning and adapting to new data. This can include adjustments to machine settings, production schedules, and maintenance plans to optimize efficiency and reduce operational costs.

18.2 K-Armed Bandits

To introduce RL learning, consider the k-armed bandit problem. It is named after the nickname of early slot machines (Figure 18.3). The RL agent is faced with k such slot machines that give different stochastic rewards. The rewards given by each of the k bandits are initially unknown to the agent. The goal of the agent is to find a policy of which bandit to play in order to maximize the return, that is, the sum of future rewards.

The k-armed bandit problem is a very simple RL problem because it is *stateless*. That is, the agent is only ever in one state and the state does not change. This means that the action value function depends only on the action, not the state-action pair.

Formally, there are k possible actions A_t at time t with stochastic reward R_t . The action value for each action can be defined as the average reward for that action:

$$Q_t(a) = \frac{\sum_{i=1}^{t-1} R_i \times \mathbb{1}_a}{\sum_{i=1}^{t-1} \mathbb{1}_a} \quad (\text{average reward})$$

Here $\mathbb{1}_a$ is 1 when action a has been taken and 0 when another action has been taken.

A suitable policy that balances exploitation of existing knowledge and exploration for gathering new knowledge is the ϵ -greedy policy. An ϵ -greedy policy is one that with probability ϵ takes a random action and with probability $1 - \epsilon$ takes the optimal action:

$$A_t = \operatorname{argmax}_a Q_t(a)$$

An incremental implementation of the action value function simply updates the running average when a new reward is received, as follows:

$$Q_{t+1}(a) = Q_t(a) + \frac{1}{t} [R_t(a) - Q_t(a)] \quad (18.2)$$

Note how the form of Equations 18.1 and 18.2 is similar. They represent different cases of the general *update rule for estimates*:

$$\text{NewEstimate} \leftarrow \text{OldEstimate} + \text{StepSize} [\text{Target} - \text{OldEstimate}]$$

Where $[\text{Target} - \text{OldEstimate}]$ is the *error* in the estimate.

A complete k-armed bandit algorithm is shown in pseudocode in Figure 18.4. The corresponding implementation in Python is straightforward¹. The following code block defines a class `k_bandit_agent` that represents an agent. Initialization specifies the number of bandits k in the environment, the parameter ϵ for the ϵ -greedy policy and the initial value of the action value function, which may be 0 as in the pseudocode in Figure 18.4. The method `determine_action` is simply the ϵ -greedy policy. The `train` method for each step determines the action to take, then takes that action in the environment and receives a reward. The agent then updates the action value function as in Equation 18.2.

¹Complete implementation is available at <https://github.com/jevermann/busi4720-rl/blob/main/bandits.py>

Initialize, for $a = 1$ to k :

$$Q(a) \leftarrow 0$$

$$N(a) \leftarrow 0$$

Loop forever:

$$A \leftarrow \begin{cases} \operatorname{argmax}_a Q(a) & \text{with probability } 1 - \epsilon \\ \text{a random action} & \text{with probability } \epsilon \end{cases}$$

$$R \leftarrow \text{bandit}(A)$$

$$N(A) \leftarrow N(A) + 1$$

$$Q(A) \leftarrow Q(A) + \frac{1}{N(A)} [R - Q(A)]$$

Figure 18.4: A simple bandit algorithm (Source: SB)

```
class k_bandit_agent:
    def __init__(self, k, epsilon, initial_value):
        self.k = k
        self.epsilon = epsilon
        self.env = k_bandit_env(k)

        self.Q = [initial_value] * self.k
        self.N = [.0] * self.k

    def determine_action(self):
        if random.uniform(0,1) < self.epsilon:
            # explore
            action = random.randint(0, self.k-1)
        else:
            # exploit
            action = self.Q.index(max(self.Q))
        return action

    def train(self, steps):
        rewards = []
        for step in range(steps):
            action = self.determine_action()
            reward = self.env.step(action)
            self.N[action] += 1
            self.Q[action] = (reward-self.Q[action])/self.N[action]
            rewards.append(reward)
        return rewards
```

A corresponding environment for the agent to act in is also readily implemented in Python and shown in the following code block. The initialization of the environment randomly sets the mean rewards of each of the k bandits. Each time a bandit is played,

the `step()` method randomly determines a reward from a standard normal distribution with the mean of the k -th bandit.

```
class k_bandit_env:
    def __init__(self, k):
        self.k = k
        self.mean_rewards = []

    for i in range(self.k):
        self.mean_rewards.append(random.normalvariate(0, 1))

    def step(self, action):
        mean = self.mean_rewards[action]
        reward = random.normalvariate(mean, 1)
        return reward
```

Figure 18.5 shows a comparison of learning behaviour for agents with different parameters ϵ . The horizontal axis shows the index of 1000 steps in the environment and the vertical axis shows the reward received at each step (mean over 1000 runs of the algorithm). The agent with $\epsilon = 0$ (blue line) shows the worst learning behaviour. That agent only exploits and never explores. In other words, it never finds any better or more valuable actions to take, once it has found a good action. In contrast, the agent with $\epsilon = 0.01$ (red line) learns slower in the beginning as it explores more but after 1000 steps has achieved a better mean reward. Finally, the purple line represents an agent with $\epsilon = 0$ but whose action value function has been “optimistically” initialized with values of 5 instead of 0, that is, above the expected reward. This prevents it from assuming the first good action is the best one, which leads to high initial learning. However, with an ϵ of 0, that agent is not capable of further learning later in the sequence of actions taken.

18.3 Markov Decision Processes and Dynamic Programming

This section introduces the concept of *Markov decision processes*, that is, a sequence of decisions or actions and states that has the Markov property: the reward and next state depend only on the current state and action, not on the state history.

One can think of RL learning as a Markov decision process. Figure 18.6 shows the RL agent and the environment it is situated in. The environment is at time t in state S_t . The agent takes action A_t in the environment and receives reward R_t . As a result, the environment’s state changes to the new state S_{t+1} . This leads to the concept of a *trajectory* as a sequence of states, actions, and rewards:

$$S_0, A_0, R_1, S_1, A_1, R_2, \dots$$

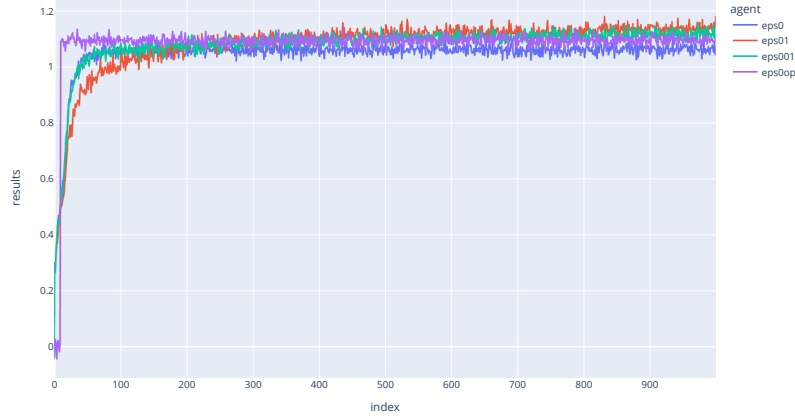
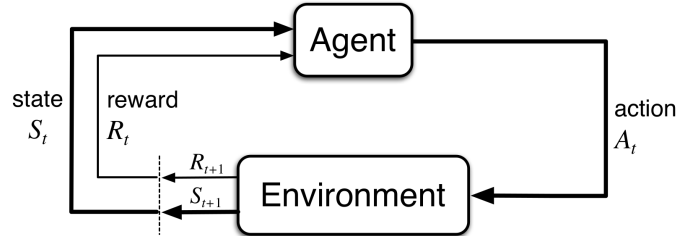


Figure 18.5: Learning performance for k-armed bandit agents for different ϵ and initial action-values



Source: SB Figure 3.1

Figure 18.6: RL agent and environment

18.3.1 Definitions

The behaviour of the environment is stochastic and can be described through the "p-function" which expresses the state transition and reward probabilities. This is known as the *dynamics* of the environment:

$$p(s', r|s, a) = \Pr\{S_t = s', R_t = r | S_{t-1} = s, A_{t-1} = a\} \quad (18.3)$$

The *return* is formally defined as the possibly discounted sum of future rewards:

$$\begin{aligned} G_t &= R_{t+1} + \gamma R_{t+2} + \gamma^2 R_{t+3} + \dots = \sum_{k=0}^{\infty} \gamma^k R_{t+k+1} \\ &= R_{t+1} + \gamma G_{t+1} \end{aligned} \quad (18.4)$$

The *state value function* of state s under a policy π is defined as the expected value of the return in that state where \mathbb{E}_π is the expectation when acting according to policy π :

$$\begin{aligned} v_\pi(s) &= \mathbb{E}_\pi[G_t | S_t = s] \\ &= \mathbb{E}_\pi \left[\sum_{k=0}^{\infty} \gamma^k R_{t+k+1} | S_t = s \right] \end{aligned} \quad (18.5)$$

Similarly, the *action value function* of state s and action a for policy π is defined as the expected return of being in state s and taking action a :

$$\begin{aligned} q_\pi(s, a) &= \mathbb{E}_\pi[G_t | S_t = s, A_t = a] \\ &= \mathbb{E}_\pi \left[\sum_{k=0}^{\infty} \gamma^k R_{t+k+1} | S_t = s, A_t = a \right] \end{aligned} \quad (18.6)$$

18.3.2 Bellman Equations and Iterative Policy Evaluation

Starting with Equation 18.5 and substituting Equation 18.4 into it yields:

$$\begin{aligned} v_\pi(s) &= \mathbb{E}_\pi[G_t | S_t = s] \\ &= \mathbb{E}_\pi[R_{t+1} + \gamma G_{t+1} | S_t = s] \end{aligned}$$

An expectation is the sum of values weighted by their probability. Summing over all possible combinations of action a , next state s' and rewards r and using the dynamics of the environment (Equation 18.3) and the stochastic policy $\pi(a|s)$ for probabilities of taking action a in state s , then yields:

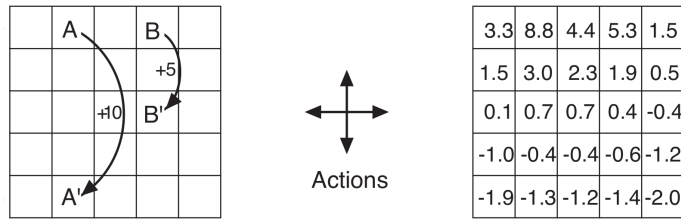
$$v_\pi(s) = \sum_a \sum_{s'} \sum_r [r + \gamma \mathbb{E}_\pi[G_{t+1} | S_{t+1} = s']] p(s', r | s, a) \pi(a | s)$$

Note that the expectation of G_{t+1} is not replaced by this sum and remains an expectation. Rearranging this slightly:

$$v_\pi(s) = \sum_a \pi(a | s) \sum_{s'} \sum_r p(s', r | s, a) [r + \gamma \mathbb{E}_\pi[G_{t+1} | S_{t+1} = s']]$$

Recognizing that the expectation in the final term on the right is just the state value function (Equation 18.5) for state s' yields:

$$v_\pi(s) = \sum_a \pi(a | s) \sum_{s', r} p(s', r | s, a) [r + \gamma v_\pi(s')] \quad \text{for all } s \in \mathcal{S} \quad (18.7)$$



Source: SB Figure 3.2

Figure 18.7: Gridworld example and optimal state value function

Equation 18.7 is called the *Bellman equation* for the state value function. A similar equation can be derived for the action value function, beginning with Equation 18.6 and following the same steps.

To illustrate the concept of the state value function, consider the gridworld example in Figure 18.7. An agent is placed on the grid in the left panel of the figure. The agent can take four possible actions; it can move up, down, left or right. Moving off the grid, for example taking action "up" when in the top row, results in a reward of -1 and the state is unchanged. All other actions yield a reward of 0 with state changes as indicated by the action, except when states A or B are reached. When reaching state A , the agent receives a reward of $+10$ and the next state is A' . When reaching state B , the agent receives a reward of $+5$ and the next state is B' .

The policy π in this example is a random policy; independent of its state, the agent takes each action with equal probability. The discount rate is set at $\gamma = 0.9$, favouring immediate rewards over future ones.

The state values for this problem under the random policy are shown in the right panel of Figure 18.7. It is clear that being in states A and B is most valuable, as the immediate reward is great. However, the state values are not $+10$ or $+5$ because the agent is likely to incur some negative future rewards. In particular, the state values of the edge states are low or negative because of the probability of falling off the world and incurring a reward of -1 .

The Bellman equations (Equation 18.7, and its equivalent for the action value function) express the fact that the value of a state (or state-action pair) is a function of the values of all other states (or state-action pairs). This suggests an intuitive way to compute the state values iteratively. Beginning with random values for each state, calculate updated values for all states using Equation 18.7. Then, consider the updated values as the current values, and calculate updated values based on these². Iterate like this until the values do not change any more. It can be proven that this procedure converges to the correct solution and terminates.

This process is called *iterative policy evaluation*. Figure 18.8 shows this in pseudocode

²In fact, it is not even necessary to wait until all states have been iterated over and updated before using updated state values as current ones.

```

Loop:
     $\Delta \leftarrow 0$ 
    Loop for each  $s \in \mathcal{S}$  :
         $v \leftarrow V(s)$ 
         $V(s) \leftarrow \sum_a \pi(a|s) \sum_{s',r} p(s',r|s,a)[r + \gamma V(s')]$ 
         $\Delta \leftarrow \max(\Delta, |v - V(s)|)$ 
    until  $\Delta < \theta$ 

```

Figure 18.8: Iterative Policy Evaluation (Source: SB)

and the following Python code block shows the straightforward implementation, beginning with initial values of 0 for each state.

```

# Define actions for gridworld
A = list(range(0,4))
# Initialize value function V
V = dict()
for state in States:
    V[state] = 0
# Initialize random policy pi
pi = dict()
for state in States:
    pi[state] = random.choice(A)

def evaluate_policy():
    while True:
        Delta = 0
        for s in States:
            v = V[s]
            V[s] = exp_reward(s, pi[s])
            Delta = max(Delta, abs(v - V[s]))
        print(Delta)
        if Delta < theta:
            break

```

18.3.3 Bellman Optimality and Iterative Policy Improvement

Maximizing the state value function v or action value function q is finding an optimal policy π , that is, that policy that when following it yields the maximum state value or action value:

$$v_*(s) = \max_{\pi} v_{\pi}(s)$$

$$q_*(s, a) = \max_{\pi} q_{\pi}(s, a)$$

Intuitively, the value of a state under an optimal policy π_* is equal to the expected return for the best action from that state:

$$v_*(s) = \max_{a \in \mathcal{A}(s)} q_{\pi_*}(s, a)$$

Substituting the definition of the action value function (Equation 18.6):

$$= \max_a \mathbb{E}_{\pi_*}[G_t | S_t = s, A_t = a]$$

Substituting the recursive definition of the return (Equation 18.4):

$$= \max_a \mathbb{E}_{\pi_*}[R_{t+1} + \gamma G_{t+1} | S_t = s, A_t = a]$$

Noting that the expected future return is just the value of the next state, that is, using Equation 18.5:

$$= \max_a \mathbb{E}[R_{t+1} + \gamma v_*(S_{t+1}) | S_t = s, A_t = a]$$

The expectation is the sum over all following states s' and rewards r weighted by their probabilities. Using the dynamics of the environment (Equation 18.3) that describe the probabilities yields:

$$v_*(s) = \max_a \sum_{s', r} p(s', r | s, a) [r + \gamma v_*(s')]$$

Similarly, the action value under an optimal policy π^* can be derived as:

$$\begin{aligned} q_*(s, a) &= \mathbb{E} \left[R_{t+1} + \gamma \max_{a'} q_*(S_{t+1}, a') | S_t = s, A_t = a \right] \\ &= \sum_{s', r} p(s', r | s, a) [r + \gamma \max_{a'} q_*(s', a')] \end{aligned}$$

These final expressions are known as the *Bellman optimality* equations for the state and action value functions.

A simple, intuitive way of finding the optimal policy is to find the optimal state value function. The optimal policy is then to take that action that will yield the best following state. However, changing the policy will change the state value function. This intuitive procedure shows that state value function calculation and policy updates should happen alternately, until the policy no longer changes.

Because the state value computation was already demonstrated using iterative policy evaluation above, the procedure for *iterative policy improvement* is simple, expressed in Figure 18.9. The following Python code block illustrates the straightforward implementation:

```

Loop:
    stable ← true
    For each  $s \in \mathcal{S}$  :
         $old\_action \leftarrow \pi(s)$ 
         $\pi(s) \leftarrow \text{argmax}_a \sum_{s',r} p(s',r|s,a)[r + \gamma V(s')]$ 
        If  $old\_action \neq \pi(s)$  then  $stable \leftarrow false$ 
    If  $stable$  then
        return  $V \approx v_*$  and  $\pi \approx \pi_*$ 
    else
        go to policy evaluation

```

Figure 18.9: Iterative Policy Improvement (Source: SB)

```

def improve_policy():
    stable = True
    for s in States:
        old_action = pi[s]
        max_r = -math.inf
        max_a = None
        for action in Actions:
            r = exp_reward(s, action)
            if r > max_r:
                max_r = r
                max_a = action
        pi[s] = max_a
        if old_action != pi[s]:
            stable = False
    return stable

```

Putting both functions, `evaluate_policy()` and `improve_policy()`, together in an iteration will yield the optimal policy:

```

stable = False
while not stable:
    evaluate_policy()
    stable = improve_policy()

print("Optimal Policy:")
print(pi)

```

The procedures of *iterative policy evaluation* and *iterative policy improvement* are an example of the more general approach to optimization called *dynamic programming*.

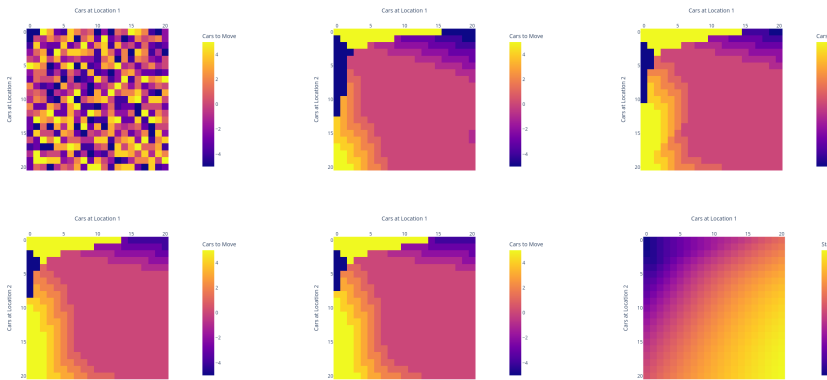


Figure 18.10: Iterative policy improvement example (exercise 4.2 in SB). Policies and final state value function.

Consider the example of "Jack's Car Rental" (example 4.2 in SB). Jack rents cars at 2 locations. Each location can store 20 cars. The number of daily rental requests and rental returns are Poisson distributed. Jack can move a maximum of 5 cars between the two locations overnight. Each move incurs a reward (cost) of -2 and each satisfied rental request receives a reward of $+10$. Jack is looking for the optimal policy that specifies how many cars to move from location 1 to location 2 every night.

The states in this problem are defined as the number of cars in location 1 and 2, for a total of $20 \times 20 = 400$ possible states and 2 actions, defined in the following Python code block³:

```
States = []
for cars1 in range(21):
    for cars2 in range(21):
        States.append((cars1, cars2))

Actions = range(-5, 5+1)
```

Figure 18.10 shows the policies after each iteration of the iterative policy improvement algorithm in Figure 18.9, beginning with the random policy in the top left panel of Figure 18.10. Positive values for the policy indicate cars to move from location 2 to location 1, negative values indicate cars to move from location 1 to location 2. The policy improvement converges to the optimal policy after 4 iterations, with the final state value function shown in the bottom right panel of Figure 18.10.

³Complete implementation available at <https://github.com/jevermann/busi4720-rl/blob/main/jacks.py>

```

Input: a policy  $\pi$  to be evaluated
Initialize:
 $V(s) \in \mathbb{R}$ , arbitrarily, for all  $s \in \mathcal{S}$ 
 $Returns(s) \leftarrow$  an empty list, for all  $s \in \mathcal{S}$ 
Loop forever (for each episode):
    Generate an episode following  $\pi : S_0, A_0, R_1, S_1, A_1, R_2, \dots, S_{T-1}, A_{T-1}, R_T$ 
     $G \leftarrow 0$ 
    Loop for each step of episode,  $t = T - 1, T - 2, \dots, 0$  :
         $G \leftarrow \gamma G + R_{t+1}$ 
        Unless  $S_t$  appears in  $S_0, S_1, \dots, S_{t-1}$  :
            Append  $G$  to  $Returns(S_t)$ 
             $V(S_t) \leftarrow \text{average}(Returns(S_t))$ 

```

Figure 18.11: First-visit MC prediction (Source: SB)

18.4 Monte Carlo (MC) Learning

The previous section illustrates how optimal policies and their state value and action value function can be computed *under the assumption that the dynamics of the environment, that is Equation 18.3, are known*. In practice, this is not the case — the $p(s', r|s, a)$ are unknown, there is no model of the environment. The RL agent must learn V and Q from *experience*, that is, it must act in an environment and generate trajectories (sequences of states, actions, and rewards).

This section assumes *episodic tasks*, that is problems with a terminal state, a finite trajectory, and finite returns. The agent acts in an environment according to a policy π until it arrives in a terminal state and the episode ends. At that point, the entire sequence of states, actions, and rewards is known and state value functions can be estimated or approximated. Recall that the value of a state is the expected return, that is the expected sum of discounted future rewards. State values can then be approximated as the average of the returns in that state.

A problem arises because the agent can be in the same state multiple times before the episode terminates. Different assumptions can be made. For example, one can assume that it is the first visit of a state that is most influential in determining the outcome of the episode and therefore the state value should be updated with the return at the time of the first visit of a state. Alternatively, one could assume that the last visit of a state is most important, and the value function is updated with the returns at the last time that the state is visited. Yet another alternative is to update the value function for all visits of a state. Figure 18.11 shows the pseudocode for this process when the state value function is updated only for the first visit of a state, known as *First-Visit Monte Carlo Prediction*.

```

Initialize for all  $s \in \mathcal{S}, a \in \mathcal{A}$ 
 $\pi(s) \in \mathcal{A}(s)$  (arbitrarily)
 $Q(s, a) \in \mathbb{R}$  (arbitrarily)
 $Returns(s, a) \leftarrow$  empty list
Loop forever (for each episode):
    Choose  $S_0 \in \mathcal{S}, A_0 \in \mathcal{A}(S_0)$  randomly
    Generate an episode following  $\pi : S_0, A_0, R_1, S_1, A_1, R_2, \dots, S_{T-1}, A_{T-1}, R_T$ 
     $G \leftarrow 0$ 
    Loop for each step of episode,  $t = T - 1, T - 2, \dots, 0$  :
         $G \leftarrow \gamma G + R_{t+1}$ 
        Unless the pair  $S_t, A_t$  appears in  $S_0, A_0, S_1, A_1, \dots, S_{t-1}, A_{t-1}$  :
            Append  $G$  to  $Returns(S_t, A_t)$ 
             $Q(S_t, A_t) \leftarrow \text{average}(Returns(S_t, A_t))$ 
             $\pi(S_t) \leftarrow \underset{a}{\text{argmax}} Q(S_t, a)$ 

```

Figure 18.12: First visit MC control with exploring starts (Source: SB)

Note the computation of the return G backwards from the end of the episode. The line "unless S_t appears in S_0, S_1, \dots, S_{t-1} " ensures the first-visit property and the final two lines approximate the state value by the average of the returns for that state.

Monte Carlo Control

While Monte Carlo prediction is useful in approximating the state value function, obtaining the optimal policy is done with *Monte Carlo control*. Instead of approximating $V(S)$ from the returns of an episode, MC control approximates $Q(S, A)$. Figure 18.12 shows the first-visit MC control algorithm in pseudocode. It is very similar to the first-visit MC prediction algorithm in Figure 18.11. Consider the final three lines: The main change is that returns are assigned not to states, but to pairs of states and actions. The action value function for a state-action pair is approximated as the average over all the returns assigned to it. The optimal policy in some state is that action for which the action value is maximal.

There is one additional difference between Figures 18.11 and 18.12 in that the initial states and actions of each episode are chosen randomly. This is necessary because the policy π in Figure 18.12 is deterministic and greedy. Without any random influence, there would be no exploration. Forcing episodes to begin with random states and actions is called *exploring starts* and it ensures that every state-action pair is visited at least once (assuming sufficiently many episodes are generated).

As an example of an episodic RL problem that can be usefully learned with MC meth-

ods, consider the game of Blackjack (example 5.3 in SB)⁴. In this game, cards have values A, 2, 3, ..., 10 where A is the ace and 10 includes face cards. An ace can count as 1 point or 11 points. An ace that is counted as 11 is called a "usable ace". The dealer's initial card is shown. The player can take two possible actions: take another card ("hit") or do not take a card ("stick"). Once the player sticks, the dealer takes cards. The dealer sticks on a sum of 17 or more. When the player's or the dealer's sum of cards is over 21 they are "bust", that is, they lose the game. The dealer's policy is deterministic, they stick on 17 points or more.

For Blackjack, the states are defined as a combination of the player's current sum of cards, the initial card the dealer is showing, and whether the player has a usable ace (one that can be converted from an 11 to a 1). Actions are to hit or stick. The following Python code block shows how this can be readily implemented:

```
gamma = 1.0
# Define states
States = []
for ace in [0,1]:
    for dealer_showing in range(1,11):
        for hand_sum in range(12, 22):
            States.append((ace,dealer_showing,hand_sum))

# Define actions
Actions = (0, 1)
```

Initially, the policy is a random policy, action value functions are 0 for all state-action pairs, and the list of returns for each state-action pair is empty:

```
# Initialize policy
pi = dict()
for s in States:
    pi[s] = random.randint(0,1)

# Initialize action value function
Q = dict()
for s in States:
    for a in Actions:
        Q[(s, a)] = 0

# Initialize returns
Returns = dict()
for s in States:
    for a in Actions:
        Returns[(s, a)] = []
```

The following code block shows the generation of an episode under policy π from initial state s_0 and with initial action a_0 . The function $\text{step}()$ calls the environment.

⁴A complete implementation is available at https://github.com/jevermann/busi4720-rl/blob/main/blackjack_es.py

It includes the dealer drawing cards after the player uses the "stick" action in a state.

```
def generate_episode(pi, s0, a0):
    terminal = False
    s = s0
    a = a0
    states = [s0]
    actions = [a0]
    rewards = [math.nan]
    while terminal is False:
        sprime, r, terminal = step(s, a)
        rewards.append(r)
        if not terminal:
            aprime = pi[sprime]
            states.append(sprime)
            actions.append(aprime)
            s = sprime
            a = aprime

    return states, actions, rewards, len(rewards)
```

The core of MC control is implemented in the following Python code block, which is very much analogous to the pseudocode in Figure 18.12. Every episode begins in a random state and with a random action. An episode is generated and the sequence of states, actions, returns, and the length of the episode T are returned. Returns are computed from the end of the episode and assigned to the first-visit of a state-action pair. The policy is updated based on the new approximate value of the Q function.

```
# Learn the Q function
for e in range(0, 1000000+1):
    s0 = random.choice(States)
    pi0 = random.choice(Actions)
    S, A, R, T = generate_episode(pi, s0, pi0)
    G = 0
    for t in reversed(range(0, T-1)):
        G = gamma*G + R[t+1]
        if (t == 0) or ((S[t], A[t]) not in zip(S[0:t-1], A[0:t-1])):
            Returns[(S[t], A[t])].append(G)
            Q[(S[t], A[t])] = mean>Returns[(S[t], A[t])])
            if Q[(S[t], 1)] > Q[(S[t], 0)]:
                pi[S[t]] = 1
            else:
                pi[S[t]] = 0
```

Figure 18.13 shows the resulting policy (left panels) and state value function (right panels) after 1,000,000 learning episodes. The top two plots in Figure 18.13 are with a usable ace, the bottom two plots without a usable ace. An action of 1 means to hit, and action 0 is to stand.

For the game of Blackjack, exploring starts with a deterministic policy are a good way to ensure exploration. However, exploring starts are not always possible or realistic.

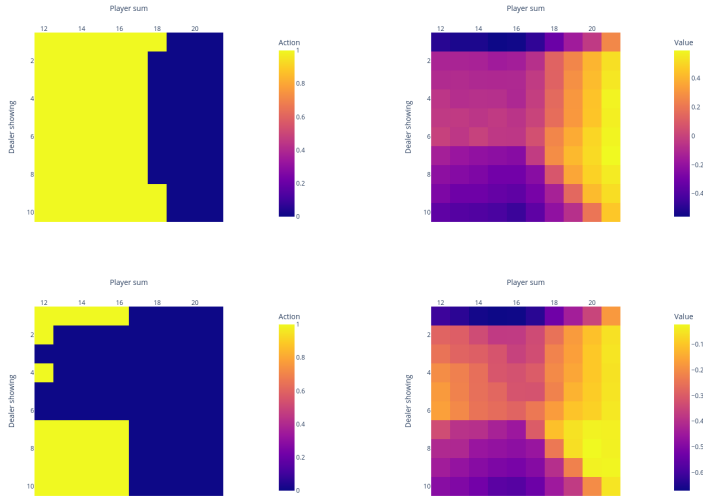


Figure 18.13: Policies and state value function for the Blackjack example after 1,000,000 episodes. Usable ace on top, no usable ace at bottom

Instead of using a greedy policy, in those cases an ϵ -soft policy can be used to ensure exploration. The policy π now represents not the action to be taken (greedily, deterministically) but the probability with which each action should be chosen. Consequently, π is updated with ϵ -soft probabilities, as shown in the following Python code fragment⁵:

```

Q[(S[t], A[t])] = mean>Returns[(S[t], A[t])])
# Optimal policy (for two actions)
A_star = 1 if Q[(S[t], 1)] > Q[(S[t], 0)] else 0

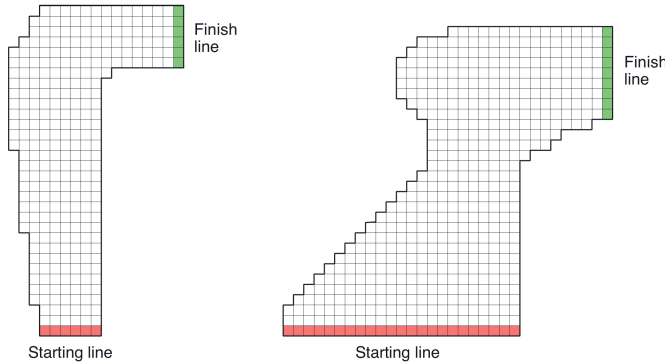
for a in Actions:
    if a == A_star:
        pi[(S[t], a)] = 1-epsilon+epsilon/len(Actions)
    else:
        pi[(S[t], a)] = epsilon/len(Actions)

```

An example where exploring starts are not realistic is the Racetrack problem (exercise 5.12 in SB). Cars have to follow the right curve of a racetrack, from the starting line to the finish line, as shown in two examples in Figure 18.14.

The states are defined by the car's position and velocity on the race track, each in two dimensions, horizontally and vertically. The possible actions are to accelerate, coast, or brake in each direction. The rewards are -1 for each step taken and +1 for crossing the finish line, when the episode terminates. When a car moves off the racetrack, it is reset

⁵ An implementation of the Blackjack problem with an ϵ -soft policy is available at https://github.com/jevermann/busy4720-rl/blob/main/blackjack_eps.py.



Source: SB Figure 5.5

Figure 18.14: Racetrack example

to a random position on the starting line and the episode continues. The environment is stochastic: With a small probability, the actions of the agents are ignored, that is, they have no effect.

The following Python code block defines the actions, the action value function and the ϵ -soft policy that samples from the probability distribution over actions in a state⁶.

```
Actions = []
for y in range(-1, 2):
    for x in range(-1, 2):
        Actions.append((y,x))

Q = dict()
def getQ(s, a):
    if (s, a) not in Q:
        return 0
    else:
        return Q[(s, a)]

pi = dict()
def get_action(s):
    weights = []
    for a in Actions:
        if (s, a) in pi:
            weights.append(pi[(s, a)])
    return random.choices(Actions, weights=weights)[0]
```

The following two Python functions manage the returns for each state-action pair:

⁶Complete implementation is available at <https://github.com/jevermann/busi4720-rl/blob/main/racetrack.py>.

```

Returns = dict()
def getReturns(s, a):
    if (s, a) not in Returns:
        return []
    else:
        return Returns[(s, a)]
def appendReturn(s, a, r):
    if (s, a) not in Returns:
        Returns[(s, a)] = [r]
    else:
        Returns[(s, a)].append(r)

```

The following Python code block represents the core MC control algorithm. Note that the policy π now represents the probabilities of taking an action in a state and is updated accordingly. The remainder of the code is largely unchanged from the Blackjack example, except that the Racetrack example does not need to use exploring starts, that is, random starting states and actions.

```

for e in range(0, 10000+1):
    S, A, R, T = env.generate_episode()
    G = 0
    for t in reversed(range(0, T-1)):
        G = gamma*G + R[t+1]
        if (t == 0) or ((S[t], A[t]) not in zip(S[0:t-1], A[0:t-1])):
            appendReturn(S[t], A[t], G)
            Q[(S[t], A[t])] = mean(getReturns(S[t], A[t]))
            A_star = argmaxQ(S[t])
            for a in Actions:
                if a == A_star:
                    pi[(S[t], a)] = 1 - eps + eps / len(Actions)
                else:
                    pi[(S[t], a)] = eps / len(Actions)

```

Figure 18.15 shows visualizations of the trajectory of a car (green line) on the racetrack after 0, 100, 200, and 10,000 episodes. It is clear from these trajectories that the number of steps is reduced as training progresses.

18.5 Off-Policy MC Learning

In the MC methods described above, the policy used to generate behaviour (“*behaviour policy*”), and the policy that is learned (“*target policy*”) are the same. However, the need to keep exploring, that is, to behave sub-optimally, when a better, greedy policy could be available, means that the agent’s performance is reduced. *Off-policy learning* is motivated by the need for efficiency and flexibility in learning optimal policies. The main motivation is the ability to learn about the optimal policy independently of the agent’s actions. This is useful in environments where exploring all actions is either potentially damaging or costly.

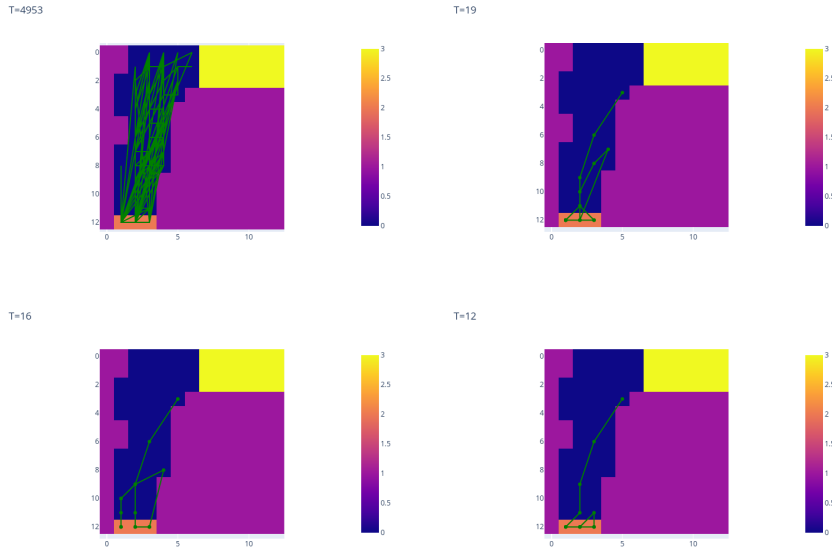


Figure 18.15: Racetrack trajectory after 0, 100, 200, and 10000 learning episodes:

In on-policy methods, the agent learns from the actions it evaluates and takes, meaning the behaviour policy is the same as the target policy. In contrast, off-policy methods allow the agent to learn a policy different from the one it follows. This is beneficial as it permits the use of historical data from other policies, or data generated from exploratory or less optimal policies, to improve a potentially different and more optimal target policy.

One important requirement in off-policy learning is that the behaviour policy must *cover* the target policy. That is, all behaviour possible under the target policy must be (eventually) generated by the behaviour policy. This requirement is intuitive, as the target policy cannot learn about behaviour, that is, the effects of actions in states, that is never encountered in a generated episode.

Figure 18.16 shows pseudocode for off-policy MC control. In this learning method, the behaviour policy b is typically an ϵ -soft policy, as presented above, that allows exploratory behaviour. The target policy π is a deterministic, greedy policy, as can be seen in the third-to-last line of Figure 18.16.

The following Python code blocks show how off-policy MC control can be implemented. The first block defines the two policies, the ϵ -soft behaviour policy b and the greedy deterministic policy π .

```

Initialize for all  $s \in \mathcal{S}, a \in \mathcal{A}(s)$  :
 $Q(s, a) \in \mathbb{R}$  (arbitrarily)
 $C(s, a) \leftarrow 0$ 
 $\pi(s) \leftarrow \operatorname{argmax}_a Q(s, a)$ 
Loop forever (for each episode):
    Generate an episode following  $b : S_0, A_0, R_1, S_1, A_1, R_2, \dots, S_{T-1}, A_{T-1}, R_T$ 
     $G \leftarrow 0; W \leftarrow 1$ 
    Loop for each step of episode,  $t = T - 1, T - 2, \dots, 0$  :
         $G \leftarrow \gamma G + R_{t+1}$ 
         $C(S_t, A_t) \leftarrow C(S_t, A_t) + W$ 
         $Q(S_t, A_t) \leftarrow Q(S_t, A_t) + \frac{W}{C(S_t, A_t)} [G - Q(S_t, A_t)]$ 
         $\pi(S_t) \leftarrow \operatorname{argmax}_a Q(S_t, a)$ 
        If  $A_t \neq \pi(S_t)$  then proceed to next episode
         $W \leftarrow W/b(A_t|S_t)$ 

```

Figure 18.16: Off-Policy MC Control (Source: SB)

```

def b(s):
    weights = []
    for a in Actions:
        if (s, a) in Q:
            weights.append(math.exp(Q[(s, a)]))
        else:
            weights.append(0)
    if len(weights) == 0 or sum(weights) == 0:
        return random.choice(Actions)
    else:
        return random.choices(Actions, weights)[0]

def pi(s):
    a = argmaxQ(s)
    if a is None:
        return random.choice(Actions)
    else:
        return a

```

Note that the last line of Figure 18.16 makes reference to the probabilities of action A_t in state S_t under the behaviour policy b . This probability is computed by the following Python code block:

```
def bprob(a, s):
    if (s, a) not in Q:
        return 1
    weights = []
    for aa in Actions:
        if (s, aa) in Q:
            weights.append(math.exp(Q[(s, aa)]))
    if len(weights) == 0 or sum(weights) == 0:
        return 1
    else:
        return math.exp(Q[(s, a)]) / sum(weights)
```

With these definitions, the learning algorithm is a straightforward implementation of the pseudocode in Figure 18.16⁷:

```
for e in range(0, 10000+1):
    S, A, R, T = env.generate_episode_b()
    G = 0
    W = 1
    for t in reversed(range(0, T-1)):
        G = gamma*G + R[t+1]
        C[(S[t], A[t])] = getC(S[t], A[t]) + W
        Q[(S[t], A[t])] = getQ(S[t], A[t]) + \
            W/getC(S[t], A[t]) * (G-getQ(S[t], A[t]))
        if A[t] != pi(S[t]):
            break
        else:
            W = W * 1/bprob(A[t], S[t])
```

18.6 Temporal-Difference (TD) Learning

Temporal Difference (TD) learning is an RL approach that combines ideas from both Monte Carlo (MC) methods and dynamic programming. The primary motivation for temporal difference learning stems from its ability to learn predictions based on other learned predictions, a concept referred to as bootstrapping. Unlike Monte Carlo methods, which wait until the completion of an episode to update the value estimates based on actual returns, TD learning updates estimates based on estimate returns, thus not requiring the episode to terminate before updates can be made. This allows TD learning to make more frequent updates, which can accelerate learning, and to be applied in continuing (non-episodic) environments.

Recall that in MC control, the updates to the action value function use the actual return G_t at time t as the target, which can only be computed at the end of an episode:

⁷Complete implementation using the Racetrack example is available at https://github.com/jevermann/busy4720-rl/blob/main/racetrack_off_policy.py.

```

Initialize  $Q(s, a)$  for all  $s \in \mathcal{S}^+$ , arbitrarily
Loop for each episode:
    Initialize  $S$ 
    Choose  $A$  from  $S$  using policy derived from  $Q$ 
    Loop for each step of episode:
        Take action  $A$ , observe  $R, S'$ 
        Choose  $A'$  from  $S'$  using policy derived from  $Q$ 
         $Q(S, A) \leftarrow Q(S, A) + \alpha [R + \gamma Q(S', A') - Q(S, A)]$ 
         $S \leftarrow S'; A \leftarrow A'$ 
    until  $S$  is terminal

```

Figure 18.17: TD-control with the SARSA method (Source: SB)

$$Q(S_t, a) \leftarrow Q(S_t, a) + \alpha [G_t - Q(S_t, a)]$$

Substituting the recursive definition of the return (Equation 18.4) yields:

$$Q(S_t, a) \leftarrow Q(S_t, a) + \alpha [R_{t+1} + \gamma G_{t+1} - Q(S_t, a)]$$

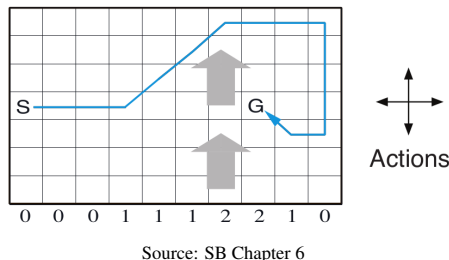
Recognizing that the expected value of G_{t+1} is approximated by the Q value of the optimal action in the next state (Equation 18.6) yields the TD update:

$$Q(S_t, a) \leftarrow Q(S_t, a) + \alpha [R_{t+1} + \gamma Q(S_{t+1}, a_{t+1}^*) - Q(S_t, a)] \quad (18.8)$$

In other words, there is no need to wait until the actual return G_t is known at the end of an episode, as TD learning uses the current approximation or estimate of G_t as the update target. This idea of using estimates to update or compute better estimates is called *bootstrapping*.

Figure 18.17 shows the pseudocode of a method called "SARSA", so-called because it uses information of the current state S , current action A , reward R , next state S' and next action A' in its update. SARSA uses the update function defined in Equation 18.8. SARSA is an on-policy method, typically using an ϵ -greedy policy based on Q to generate behaviour and ensure exploration.

To illustrate the use of SARSA, consider the "Windyworld" example environment (example 6.5 in SB), shown in Figure 18.18. An agent has to move from start state S to terminal goal state G in this gridworld. However, there is a wind on some columns of the grid that pushes the agent towards the top (indicated below the columns in Figure 18.18). Rewards are -1 for every step until termination, so that the agent is encouraged to find the path requiring the least number of actions. There are no penalties for



Source: SB Chapter 6

Figure 18.18: Windyworld example

moving off the world (the action is simply ignored) and the problem is not discounted ($\gamma = 1$).

The following python code blocks define the states, actions, and an ϵ -greedy policy π :

```
# Define states
States = []
for i in range(nrow):
    for j in range(ncol):
        States.append((i, j))

# Define actions
Actions = range(0, 4)
# Initialize Q
Q = dict()
for s in States:
    for a in Actions:
        Q[(s, a)] = random.random()

# Define pi
def pi(s):
    if random.random() < epsilon:
        return random.choice(Actions)
    else:
        return argmaxQ(s)
```

With these definitions, the implementation of SARSA is straightforward from the pseudocode in Figure 18.17⁸ and shown in the following Python code block.

⁸A complete implementation of the Windyworld example is available at https://github.com/jevermann/busy4720-rl/blob/main/windyworld_sarsa.py.

```

for e in range(0, 100):
    terminal = False
    S = windy.reset()
    A = pi(S)
    step = 0
    while terminal is False:
        Sprime, R, terminal = windy.step(A)
        Aprime = pi(Sprime)
        Q[(S,A)] = Q[(S,A)] + alpha*(R + \
            gamma * Q[(Sprime, Aprime)] - Q[(S, A)])
        S = Sprime
        A = Aprime

```

In summary, Temporal Difference learning and Monte Carlo methods differ primarily in when and how the updates to value estimates are made:

- *Update Timing*: TD learning updates values at every time step using current estimates, which means it can start learning from incomplete sequences, making it suitable for non-episodic environments. Monte Carlo methods update only at the end of each episode, using the total accumulated return from the episode.
- *Sampling vs. Bootstrapping*: Monte Carlo methods rely solely on actual returns (full sampling), and do not bootstrap. In contrast, TD methods bootstrap, using existing value estimates to update new estimates.
- *Convergence Properties*: Due to its incremental nature and frequent updates, TD learning can converge faster in practical applications than Monte Carlo methods, which require longer trajectories and may suffer from higher variance in their estimates due to the complete reliance on actual returns.

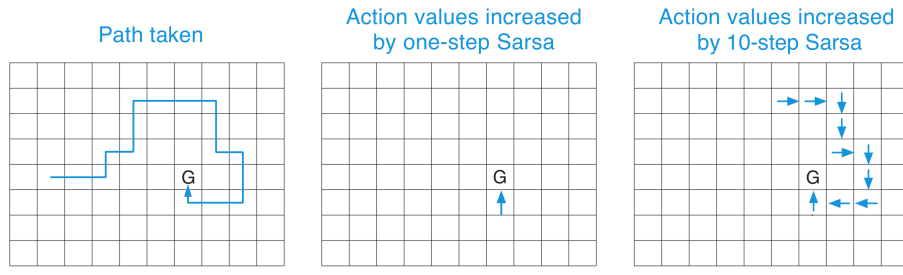
Generalizing TD Learning to N Steps

In developing the SARSA method (Equation 18.8), the recursive definition of the return (Equation 18.4) and the definition of the action value as the expected return (Equation 18.6) were applied once. Hence the SARSA update target $R_{t+1} + Q(S_{t+1}, a_{t+1})$ "looks ahead" by one step to the next reward R_{t+1} and then approximates the remaining portion of G_{t+1} by $Q(S_{t+1}, a_{t+1})$. This is therefore called the "1-step" target, and the update error δ is called the "1-step error":

$$\delta_{TD1} = R_{t+1} + Q(S_{t+1}, a_{t+1}) - Q(S_t, a)$$

This can be extended by applying Equations 18.4 and 18.6 a second time. This yields the "2-step error" that looks ahead at the next two rewards, and then approximates the remainder by $Q(S_{t+2}, a_{t+2})$:

$$\delta_{TD2} = R_{t+1} + \gamma R_{t+2} + \gamma^2 Q(S_{t+2}, a_{t+2}) - Q(S_t, a)$$



Source: SB Figure 7.4

Figure 18.19: SARSA versus n-Step TD Learning (n-step SARSA)

Calculating the 2-step error requires knowledge of two actual rewards, so the 2-step update can be performed only after two steps. Or, viewed from a different perspective, the 2-step update updates not only the value of the most recent state-action pair but the values of the two most recent state-action pairs.

This can be generalized by applying Equations 18.4 and 18.6 n -times, yielding the "n-Step TD error":

$$\delta_{TDn} = R_{t+1} + \gamma R_{t+2} + \cdots + \gamma^{n-1} R_{t+n} + \gamma^n Q(S_{t+n}, a_{t+1}) - Q(S_t, a)$$

The n -step update can only be performed after n steps have passed so that n actual rewards are available, with the remainder of G_{t+1} being approximated. As an alternative interpretation, the n -step update changes the values of the last n state-action pairs in the trajectory. Figure 18.19 shows an illustration of this for 10-step SARSA.

18.7 Off-Policy TD Learning

Temporal Difference learning can be combined with off-policy learning as well. A popular method for this is Q-learning. Q-learning updates the action values in a greedy way, that is, of a greedy policy, regardless of the action taken by the policy being followed (the behavior policy). The differences between SARSA and Q-learning are minor. Importantly, it is not necessary to explicitly encode the behaviour and target policies.

Consider the following aspects of SARSA. The next action A' that is carried out is determined using a (behaviour) policy based on Q and Q is being learned (updated) based on the actually taken action A' (target policy). This makes SARSA an on-policy method.

SARSA (on-policy):

Take action A , observe R, S'
 Choose A' from S' using policy derived from Q

$$Q(S, A) \leftarrow Q(S, A) + \alpha [R + \gamma Q(S', A') - Q(S, A)]$$

$$S \leftarrow S'; A \leftarrow A'$$

A minor change is sufficient to change on-policy SARSA to off-policy Q-learning, shown in the box below. Here, the next action A that is carried out is also determined using a (behaviour) policy based on Q , but Q is updated not based on the actually taken action A , but based on the optimal action A' , that is, the action with the maximum Q value in the following state S' . This difference means that the policy that governs behaviour ("behaviour policy") is different than the policy that is updated or learned ("target policy"), making Q-learning an off-policy method.

Q-learning (off-policy):

Choose A from S using policy derived from Q
 Take action A , observe R, S'

$$Q(S, A) \leftarrow Q(S, A) + \alpha \left[R + \gamma \max_a Q(S', A') - Q(S, A) \right]$$

$$S \leftarrow S'$$

The corresponding Python implementation for the Windyworld example is also straightforward from the above box⁹:

```
for e in range(0, 1000):
    terminal = False
    S = windy.reset()
    step = 0
    while terminal is False:
        A = pi(S)
        Sprime, R, terminal = windy.step(A)
        Q[(S,A)] = Q[(S,A)] + alpha*(R + \
            gamma * maxQ(Sprime) - Q[(S, A)])
        S = Sprime
```

Figure 18.20 shows the difference in learning behaviour between SARSA and Q-learning on the Windyworld problem. It plots the the number of required actions to achieving the goal against the number of episodes generated; off-policy Q-learning shows faster learning than SARSA.

⁹A complete implementation is available at https://github.com/jevermann/busi4720-rl/blob/main/windyworld_q_learning.py.

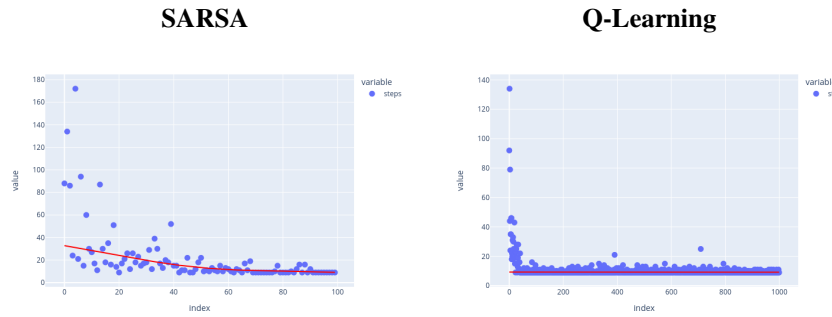


Figure 18.20: SARSA and Q-Learning Results on Windyworld

18.8 Review Questions

Introduction

1. What is reinforcement learning and how does it differ from other types of machine learning?
2. What is meant by the term "model" in the context of reinforcement learning?
3. Discuss the challenges associated with the stochastic nature of the environment in reinforcement learning.
4. Define the following terms and explain their significance in reinforcement learning:
 - Policy (π)
 - Reward (R)
 - Return (G)
 - State value function (v)
 - Action value function (q)
 - Model (p)
5. Explain the difference between a deterministic policy and a stochastic policy. Provide an example of each from any real-world scenario.
6. Describe the general process of updating the value function. Include the update rule equation and explain each component.
7. Explain the significance of the action value function in the reinforcement learning framework. How does it differ from the state value function in terms of utility and information provided to the agent?
8. Discuss how reinforcement learning can be applied to customer interaction management and financial portfolio management. What are the potential benefits and challenges?
9. Consider the manufacturing process optimization application of reinforcement learning. Describe how an RL algorithm could continuously improve the process and what metrics it might focus on.
10. Discuss the implications of the exploration-exploitation trade-off in a non-gaming business scenario. How might a company balance these two aspects effectively?

11. Reflect on the potential impacts of reinforcement learning on customer satisfaction in a service-oriented business. What are the risks and rewards?
12. What role does the discount factor γ play in calculating the return in a reinforcement learning problem? Why might one use a smaller or larger value of γ ?

Questions on K-Armed Bandits

13. Why is the k-armed bandit problem considered to be a "stateless" problem in reinforcement learning?
14. Describe the action value function $Q_t(a)$ in the k-armed bandit problem. What does it represent?
15. Explain the concept of the ϵ -greedy policy. How does it balance the exploration-exploitation trade-off?
16. Consider the incremental update formula used in the k-armed bandit problem and explain each term: $Q_{t+1}(a) = Q_t(a) + \frac{1}{t}[R_t(a) - Q_t(a)]$. How does this formula ensure that the estimate becomes more accurate over time?
17. What are the implications of setting a higher or lower ϵ value in terms of long-term gains vs. short-term exploration?
18. Discuss the rationale behind using an "optimistic" initial value for the action-value estimates. How does this approach influence the agent's behavior?

Questions on Markov Decision Processes and Dynamic Programming

19. Define a Markov Decision Process (MDP) and explain the Markov property in this context.
20. Draw a figure of an RL agent and environment interaction; describe the roles of S_t , A_t , and R_t .
21. Explain what is meant by a *trajectory* in the context of reinforcement learning.
22. Define the environment's dynamics using the *p-function* and discuss its importance in MDPs.
23. Define the *state value function* $v_\pi(s)$ and explain how it is used to evaluate a policy π .
24. Similarly, define the *action value function* $q_\pi(s, a)$ and explain its relevance in policy evaluation.
25. Detail how the Bellman equation provides a recursive way to compute the value of a state under a specific policy.
26. Describe how iterative policy evaluation can be used to approximate the state value function before performing policy improvement.
27. Explain the process of *iterative policy improvement* and how it leads to finding an optimal policy.

Questions on Monte Carlo (MC) Learning

28. Explain the fundamental concept of Monte Carlo (MC) learning in the context of reinforcement learning.
29. Define and distinguish between episodic tasks and continuous tasks in reinforcement learning.

30. Explain the difference between first-visit and every-visit MC methods. What are the implications of each approach on the learning process?
31. Describe the process of First-Visit Monte Carlo prediction as pseudocode. How does it update the state value function?
32. What is the purpose of the backward computation of the return G in the first-visit MC prediction algorithm?
33. Explain the concept of "exploring starts" in the context of MC control and why it is necessary.
34. How do Monte Carlo methods handle the trade-off between exploration and exploitation, especially in environments where exploring starts are not feasible?
35. How does the game of Blackjack illustrate the application of Monte Carlo methods in episodic RL problems?
36. Analyze the potential impact of episodic task length on the effectiveness of Monte Carlo methods. What happens as the length of episodes increases?
37. How do Monte Carlo methods adapt when the environment's dynamics change over time?
38. Reflect on how Monte Carlo methods might perform in real-world scenarios such as financial markets or automated driving systems.

Questions on Off-Policy Monte Carlo (MC) Learning

39. Explain the difference between on-policy and off-policy learning methods in the context of Monte Carlo (MC) methods.
40. How does off-policy learning address the exploration-exploitation dilemma differently than on-policy learning?
41. Discuss why it is important for the behavior policy to cover the target policy in off-policy MC learning.
42. How does the update formula in off-policy MC control differ from the one used in on-policy MC control?
43. In the pseudocode for off-policy MC control, why is the episode terminated early if the action taken does not match the action recommended by the target policy?
44. Explain the potential impacts of the choice of behavior policy on the efficiency and effectiveness of off-policy learning.
45. Discuss how off-policy learning can be applied to complex environments where safety or cost constraints limit exploration.
46. Describe a scenario in which off-policy learning would be particularly advantageous over on-policy learning.
47. How can off-policy MC control be adapted to environments where the behavior policy cannot sufficiently cover the target policy?

Questions on Temporal-Difference (TD) Learning

48. What is temporal-difference (TD) learning and how does it differ from Monte Carlo methods?
49. Explain the concept of bootstrapping in the context of TD learning.

50. Discuss the advantages of TD learning in terms of update frequency and applicability to different types of environments.
51. Describe the SARSA algorithm and explain how it uses the TD update formula.
52. How can the SARSA algorithm be adjusted to improve its performance in highly dynamic environments?
53. Discuss how TD learning methods can be adapted to continuous (non-episodic) environments. What are the implications for learning in such environments?
54. Compare the convergence properties of TD learning and Monte Carlo methods. Why might TD learning converge faster in practical applications?
55. Explain the concept of "n-step TD learning." How does it extend the basic idea of TD learning?
56. What are the implications of using different "n" values in n-step TD learning on the performance and speed of learning?
57. Explain how n-step TD learning might provide a more stable learning update compared to one-step TD updates.
58. Provide an example scenario where TD learning might significantly outperform Monte Carlo methods in terms of learning efficiency and accuracy.

Questions on Off-Policy Temporal-Difference (TD) Learning

59. Describe the main difference between the update rules of SARSA and Q-learning. How does this difference define each as either on-policy or off-policy?
60. Explain how the Q-learning update rule ensures that the learning is directed towards the optimal policy.
61. What are the implications of using the $\max_a Q(S', A')$ term in the Q-learning update rule? Discuss how this term influences the policy improvement process.
62. How does Q-learning handle the exploration-exploitation trade-off differently compared to SARSA?
63. How does the initial setting of Q-values influence the learning process and eventual performance in Q-learning? Discuss the impact of optimistic versus pessimistic initialization.

Chapter 19

Reinforcement Learning – Function Approximation

Sources and Further Reading

The material in this chapter is based on the following sources.

Richard S. Sutton and Andrew G. Barto (2018) *Reinforcement Learning – An Introduction*. 2nd edition, The MIT Press, Cambridge, MA. (SB)
<http://incompleteideas.net/book/the-book.html>

Chapters 9–13

(CC BY-NC-ND License)

The Sutton & Barto book is a standard introductory textbook on reinforcement learning and widely used. It is very approachable, but at the same time also detailed and thorough in its exposition. Its focus is on RL prior to the use of neural networks for function approximation, so up to about 2015. While it does not provide Python code itself, the pseudo-code in the book is easily implemented.

Sudharsan Ravichandiran (2020) *Deep Reinforcement Learning with Python*.
2nd edition. Packt Publishing, Birmingham, UK.
Chapters 9–11

The book by Ravichandiran is practically oriented with plenty of Python code. It discusses some of the theoretical background, but does not go into depth. It should be used after reading the Sutton & Barto chapters on function approximation and policy-based

methods.

Resources

Complete implementations of all examples in this chapter are available on the following GitHub repo:

<https://github.com/jevermann/busi4720-rl>

The project can be cloned from this URL:

<https://github.com/jevermann/busi4720-rl.git>

19.1 Introduction

In tabular RL methods the value of each state or state-action pair is represented explicitly in a table. However, as the complexity of environments grows, particularly with a high number of states or continuous state spaces, tabular methods become infeasible due to their extensive memory requirements.

To address these scalability issues, function approximation methods are employed. Function approximation techniques involve using a parameterized function to represent the value functions or the policy, rather than storing them explicitly for each state or state-action pair. In addition to addressing the scalability problem, this approach also facilitates generalization across states thereby improving learning efficiency and enabling RL to be applied to more complex and realistic problems. The types of function approximators commonly used in RL are linear functions and neural networks.

Function approximation can be applied to the state values v , the action values q and directly to the policy π :

- Approximate the state value $v(s)$ by a parameterized function $\hat{v}(s)$ with a parameter vector θ :

$$\hat{v}(s) = \hat{v}(s, \theta) \approx v_\pi(s)$$

- Approximate the action-value function $q(s, a)$ by a parameterized function $\hat{q}(s, a)$ with a parameter vector θ :

$$\hat{q}(s, a) = \hat{q}(s, a, \theta) \approx q_\pi(s, a)$$

- Approximate the policy $\pi(a, s)$ by a parameterized function $\hat{\pi}(a, s)$ with a parameter vector θ :

$$\hat{\pi}(a|s) = \hat{\pi}(a|s, \theta) \approx \pi(a|s)$$

Function approximation methods offer several advantages over traditional tabular approaches:

- *Scalability:* They can handle large or continuous state spaces efficiently.
- *Generalization:* Because changes to the parameter vector θ affect the values of multiple states or actions, function approximation methods can generalize from seen to unseen states, which is particularly useful in environments where experiencing all possible states is impractical.
- *Flexibility:* They can be adapted to different problems by choosing appropriate functions, such as linear functions or neural networks. This makes them suitable for a wide variety of problems.
- *Efficiency:* Because updates to θ affect multiple states, function approximation methods may experience improved learning and faster convergence.
- *Observability:* They can be applied to partially observable problems, as the state function need not depend on the complete state information.

Despite their advantages, function approximation methods introduce new challenges:

- *Stability and Convergence:* The use of approximators can lead to instability and divergence in some cases, particularly when combined with off-policy learning.
- *Complexity of Design:* Choosing the right features, architecture, or kernel functions requires domain knowledge and careful engineering.
- *Overfitting:* There is a risk of overfitting to the peculiarities of the sampled data, especially with highly flexible models like deep neural networks.

19.2 Value-Based Methods and Stochastic Gradient Descent

Function approximation aims to minimize the differences between the true state or action value function and the approximated function. Assuming a MSE loss, the *value error* VE can be expressed as follows:

$$\text{VE} = \sum_{s \in \mathcal{S}} \mu(s) [q_\pi(s, a) - \hat{q}(s, a, \theta)]^2$$

Stochastic gradient descent (SGD) is used to minimize this loss function, similar to the use of SGD in neural network machine learning. Refer to that chapter for a discussion of problems that can arise with SGD and different optimization methods that address these problems.

The parameters θ are iteratively updated using the gradient of the loss function. Intuitively, this process follows the steepest slope ("gradient," vector of partial derivatives) of the function to update the parameters:

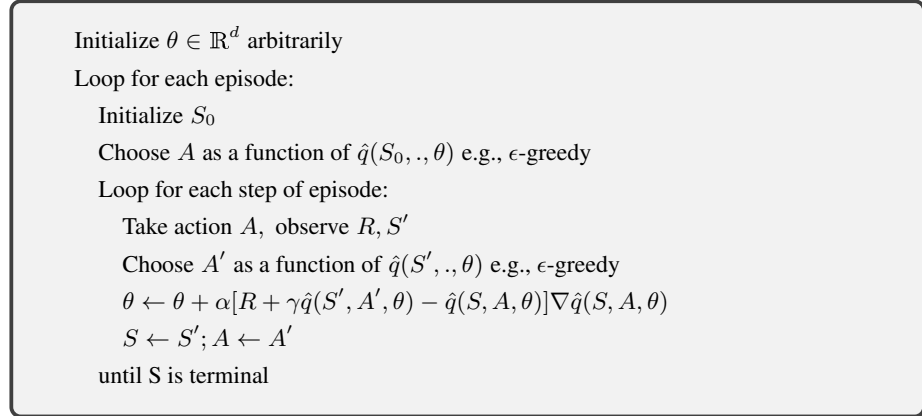


Figure 19.1: Semi-gradient SARSA (Source: SB)

$$\begin{aligned}\theta_{t+1} &= \theta_t - \frac{1}{2}\alpha\nabla [q_\pi(S_t, A_t) - \hat{q}(S_t, A_t, \theta_t)]^2 \\ &= \theta_t + \alpha[q_\pi(S_t, A_t) - \hat{q}(S_t, A_t, \theta_t)]\nabla\hat{q}(S_t, A_t, \theta_t)\end{aligned}$$

Unfortunately, the true values $q_\pi(S_t, A_t)$ are unknown. However, using the idea of bootstrapping and the definition of the action value function as an estimate of the return means that the following expression U_t can be used as an estimate of the true value $q_\pi(S_t, A_t)$:

$$U_t = R_t + \gamma\hat{q}(S_{t+1}, A_{t+1}, \theta_t) \approx q_\pi(S_t, A_t)$$

Then the parameter update becomes:

$$\begin{aligned}\theta_{t+1} &= \theta_t + \alpha[U_t - \hat{q}(S_t, A_t, \theta_t)]\nabla\hat{q}(S_t, A_t, \theta_t) \\ &= \theta_t + \alpha[R_t + \gamma\hat{q}(S_{t+1}, A_{t+1}, \theta_t) - \hat{q}(S_t, A_t, \theta_t)]\nabla\hat{q}(S_t, A_t, \theta_t)\end{aligned}\tag{19.1}$$

While tabular methods update the value of Q for a state–action pair directly, function approximation methods replace the update to Q with an update to θ . This updates the values of Q for many state–action pairs indirectly, as Q is approximated by a parameterized function.

Figure 19.1 shows how the tabular SARSA method can be readily adapted to function approximation using SGD. *The only change to the tabular SARSA algorithm is the update step.* Whereas tabular SARSA updates $q(s, a)$, semi-gradient SARSA updates the parameters θ of \hat{q} .

19.3 Deep Q Network (DQN)

While conceptually sound, simple function approximation implementations like SARSA in Figure 19.1 have some problems in practice. In particular, *instability* and *divergence* of learning arise when combining the following three elements in an RL method. These are colloquially known as the "*deadly triad*" of reinforcement learning.

- *Function approximation*: Generalizing from a state space using linear functions or neural networks.
- *Bootstrapping*: Targets include existing estimates (e.g. SARSA) rather than actual rewards only (e.g. MC methods).
- *Off-policy training*: Training on a distribution of state transitions other than that produced by the target policy.

To address these problems, RL implementations use experience replay and separate target parameters (or target networks when functions are neural networks).

Experience replay

Experience replay is a technique to break the auto-correlation between the q values of successive training batches by smoothing changes in the data distribution between mini-batches, thus making training more stable. Rather than using the generated tuple of (S, A, R, S', A') immediately in an update as in Figure 19.1, these tuples are stored in a *replay buffer*. The replay buffer is a FIFO (first-in, first-out) queue of fixed size; when it is full, older elements are removed from the front of the queue as new elements are added to the back of the queue. For every parameter update step, a sample is randomly taken from the replay buffer to fill a training batch for the SGD update step.

Target network

Working with two different sets of parameters θ_T and θ_M , one for computing the update targets $R + \gamma \hat{q}(S', A', \theta_T)$, called the "*target parameters*" and one for computing the current estimates $\hat{q}(S, A, \theta_M)$, called the "*main parameters*", has the advantage that stable update targets are provided for multiple SGD update steps. This also stabilizes training. Periodically, the target parameters are updated with the main parameters.

Because the approximation functions are typically neural networks, target parameters and main parameters are the weights and biases of two neural networks with identical architecture. Hence, one uses the terms "*target network*" and "*main network*".

Taking the two ideas of experience replay and target networks and adapting the gradient SARSA algorithm in Figure 19.1 leads directly to the DQN algorithm shown in Figure 19.2.

In practice, the state S is a function $\phi(X)$ of some raw inputs X through feature-extraction and pre-processing. To further stabilize learning, in practice the update $[y_j - \hat{q}_M(S_j, A_j, \theta_M)]$ in Figure 19.2 is clipped to $[-1, 1]$.

```

Initialize replay buffer  $D$ 
Initialize main action-value function approximation  $\hat{q}_M$  with random parameters  $\theta_M$ 
Initialize target action-value function approximation  $\hat{q}_T$  with parameters  $\theta_T = \theta_M$ 
Loop for each episode:
    Initialize  $S$ 
    For each step of the episode:
        Select action  $A$  using an  $\epsilon$ -greedy policy based on  $\hat{q}_M$ 
        Take action  $A$  and observe  $R, S_{t+1}$ 
        Store transition  $(S_t, A_t, R_t, S_{t+1})$  in  $D$ 
        Sample minibatch  $(S_j, A_j, R_j, S_{j+1})$  from  $D$ 
        Target  $y_j \leftarrow \begin{cases} r_j & \text{if } S_{j+1} \text{ is terminal} \\ r_j + \gamma \max_{A'} \hat{q}_T(S_{j+1}, A'; \theta^-) & \text{otherwise} \end{cases}$ 
         $\theta \leftarrow \theta + \alpha[y_j - \hat{q}_M(S_j, A_j, \theta_M)]\nabla \hat{q}_M(S_j, A_j, \theta_M)$ 
    Every  $C$  steps, update  $\hat{q}_T \leftarrow \hat{q}_M$  by setting  $\theta_T \leftarrow \theta_M$ 

```

Figure 19.2: DQN Algorithm (adapted from SB)

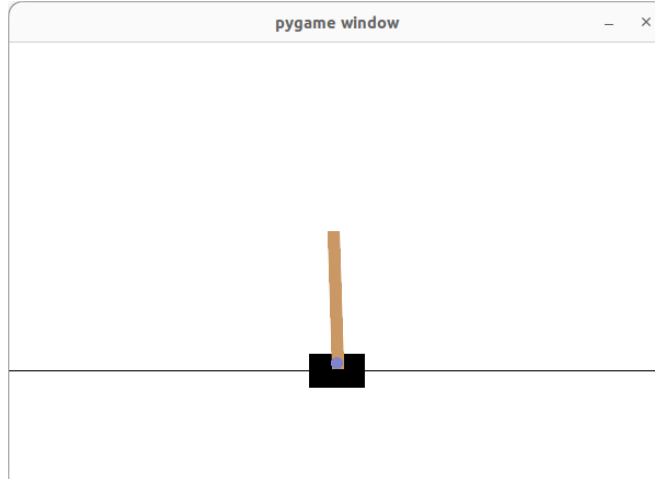


Figure 19.3: CartPole environment

Example

To illustrate a simple DQN example, consider the "Cart Pole" problem, shown in Figure 19.3. The goal is to balance an upright pole on a cart that can move left or right (but not remain still). The pole obeys a simplified physics and can tip over to the left or right.

In this particular version of the problem, the *action space* is binary, 0 pushes the cart to the left, 1 pushes the cart to the right. Every state is characterized by four *features* $x_1 \dots x_4$, the cart position ($-4.8 \leq x_1 \leq 4.8$), the cart velocity ($-\infty \leq x_2 \leq \infty$), the pole angle ($-24^\circ \leq x_3 \leq 24^\circ$), and the pole angular velocity ($-\infty \leq x_4 \leq \infty$). The *rewards* are +1 for every step taken. This means the agent has to try to prevent the pole from tipping over to get the greatest return. *Termination* of the episode occurs when the cart is moving out of range (cart position $|x_1| > 2.4$), the pole is tipping over (pole angle $|x_3| > 12^\circ$), or the episode length is greater than 200.

The DQN can be implemented using the "CartPole" environment¹. First, the required packages are loaded, the environment is created and the number of actions and number of features of a state are determined:

```
import math
import random
import keras
from keras import layers
import gymnasium as gym
import tensorflow as tf
import numpy as np
import pygame

env = gym.make("CartPole-v1", render_mode="human")

Actions = range(0, env.action_space.n)
Ssize = env.observation_space.shape[0]
```

The next Python code block defines hyperparameters for the neural network and for reinforcement learning:

```
# Neural net parameters
batch_size = 20
dropout = 0.25
activation = 'relu'

# Reinforcement learning parameters
epsilon = 0.05 # epsilon
gamma = 0.9 # discount factor
C = 5*batch_size # When to update weights

# Replay buffer D
D = collections.deque(maxlen=5000)
```

The functions \hat{q}_M and \hat{q}_T are sequential, fully-connected neural networks with a single output unit, defined in Keras. The output represents the value of \hat{q} computed by the network from its inputs. The inputs are state-action pairs, which is why

¹<https://gymnasium.farama.org>. The Farama gymnasium provides a number of reference environments for reinforcement learning.

`input_shape=(Ssize+1)` in the Python code block below which defines the main neural network:

```
# Main network, used to select actions
Q_m = keras.Sequential([
    layers.InputLayer(input_shape=(Ssize+1),
                      batch_size=batch_size,
                      dtype=tf.float32),
    layers.Dense(Ssize*4, activation=activation),
    layers.Dropout(rate=dropout),
    layers.Dense(Ssize*2, activation=activation),
    layers.Dropout(rate=dropout),
    layers.Dense(1, activation='linear')
])
Q_m.compile(loss='huber', optimizer='adam')
```

Keras provides functions that make cloning a network and getting and setting weights easy. The following Python code block creates the target network as a copy of the main network and sets its weights to those of the main network:

```
# Target network, used to compute targets
Q_t = keras.models.clone_model(Q_m)
Q_t.compile(loss='huber', optimizer='adam')
Q_t.set_weights(Q_m.get_weights())
```

Getting a value of \hat{q} for some input state-action pair is prediction from the network. The following function prepares the inputs (state features and action) as a Numpy array, adding the minibatch dimension, then selects the first prediction of the first return batch:

```
def getQ(Q, s, a):
    return Q.predict(
        np.expand_dims(np.array(s.tolist()+[a]), axis=0), \
        verbose=0)[0][0]
```

The following Python code block implements a convenient max and argmax function over all actions for the \hat{q} values of the main or target network:

```
def maxQ(Q, s, arg):
    maxq = -np.inf
    maxa = None
    for a in Actions:
        q = getQ(Q, s, a)
        if q > maxq:
            maxq = q
            maxa = a
    return maxa if arg else maxq
```

The policy π is an ϵ -greedy policy, defined in Python in the following code block. This uses the main network Q_m .

```
def pi(s, epsilon):
    if random.random() < epsilon:
        return random.choice(Actions)
    else:
        return maxQ(Q_m, s, True)
```

The update target for the DQN uses the target network Q_t and the target expression from Figure 19.2.

```
def target_DQN(Q_t, r, sprime):
    return r + gamma * maxQ(Q_t, sprime, False)
```

The following function takes a batch of entries of the experience replay buffer and creates training batches of inputs x (state features and action) and target y :

```
def training_xy(batch):
    x = np.zeros((batch_size, Ssize+1))
    y = np.zeros(batch_size)
    for i, (s, a, r, t, sprime) in enumerate(batch):
        x[i] = list(s) + [a]
        if t == 1:
            y[i] = r
        else:
            y[i]=target_DQN(Q_t, r, sprime)
    return x, y
```

The final block of Python code is the DQN algorithm, a straightforward implementation of Figure 19.2². The SGD update step is done using the Keras function `train_on_batch` that trains the network on a single batch of data.

²A complete implementation is available at https://github.com/jevermann/busi4720-rl/blob/main/DDQN_tuples.py.

```

t = 0
for episode in range(max_episodes):
    s = env.reset()[0]
    terminal = False
    while not terminal:
        a = pi(s, epsilon)
        sprime, r, terminal, _, _ = env.step(a)
        t += 1
        D.append((s, a, r, int(terminal), sprime))
        s = sprime
        if t >= batch_size:
            batch = random.sample(D, batch_size)
            x, y = training_xy(batch, ddqn=False)
            loss = Q_m.train_on_batch(x=x, y=y)

    if t % C == 0:
        Q_t.set_weights(Q_m.get_weights())

```

Double DQN

An extension to the DQN algorithm is the Double DQN (DDQN). It is based on the idea of Double-Q learning for tabular methods and uses the target network \hat{q}_T as a second Q function. This removes the upward bias from using the $\max()$ function as target estimator. The only change to be made is in the definition of the target, which, for a DDQN is:

$$\text{Target } y_j \leftarrow \begin{cases} r_j & \text{if } S_{j+1} \text{ is terminal} \\ r_j + \gamma \hat{q}_T(S_{j+1}, \arg\max_{A'} \hat{q}_M(S_{j+1}, A')) & \text{otherwise} \end{cases}$$

In Python, this is also a simple change:

```

def target_DDQN(Q_m, Q_t, a, r, sprime):
    return r + gamma * getQ(Q_t, sprime, maxQ(Q_m, sprime, False))

```

Prioritized Replay

Another extension to the basic DQN algorithm is the use of *prioritized replay*. In the DQN algorithm above, sampling from the experience replay buffer was done with uniform probability for all elements in the buffer. However, there are some elements that are more informative than others, that is, more can be learned from them than from others. In particular, these are the elements that have a large absolute TD error, that is, the elements for which $|y_j - \hat{q}_M(S_j, A_j, \theta_M)|$ is large, where y_j is either the DQN or DDQN target. Intuitively, elements that have a small prediction error are not very informative, as not much can be learned from them. When using prioritized experience

replay, the TD errors are calculated when experience tuples are added to the replay buffer. Sampling from the buffer takes the priorities into account.

Dueling DQN

The Dueling DQN is another extension of the basic DQN algorithm. It is based on the *advantage function*, which is the difference between the action value function and the state value function:

$$A(s, a) = Q(s, a) - V(s)$$

In other words, the advantage function expresses the advantage of taking action a in state s over the average action in state s that is represented by the state value function. The advantage function can be rewritten as follows:

$$Q(s, a) = V(s) + A(s, a)$$

This formulation of the advantage function suggests that the action value function can be composed of two functions. In practice, that means the computation of the action value function is done by two different neural networks, the "value stream" and the "advantage stream". Both use the same state features x as input. The advantage stream additionally receives the action a as input. In practice, the value stream and advantage stream use one or more common neural network layers, e.g. dense layers, and then separate to end in two different output nodes, one for the value function and one for the advantage function. The two outputs are then added to calculate the action value function q as follows:

$$Q(s, a, \theta, \alpha, \beta) = V(s, \theta, \beta) + \left(A(s, a, \theta, \alpha) - \frac{a}{|\mathcal{A}|} A(s, a', \theta, \alpha) \right)$$

Here, θ are shared neural-network parameters, β are parameters only for the "value-stream" neural network, and α are parameters only for the "advantage-stream" neural network.

19.4 Policy Gradient Methods

Policy gradient methods optimize the policy directly. Unlike value-based methods, which first estimate the action value function and derive a policy based on these estimates, policy gradient methods adjust the policy parameters θ directly in response to the received reward. This direct approach enables more nuanced strategies and behaviors, particularly in environments with high-dimensional or continuous action spaces.

Policy gradient methods rely on optimizing parameterized policies with respect to the expected return by gradient ascent. The policy is typically represented as

$$\pi(s, a) = \pi(s, a | \theta) = \Pr(A_t = a | S_t = s, \theta_t = \theta)$$

which defines the probability of selecting action a in state s , parameterized by θ .

The objective function in policy gradient methods is defined as:

$$J(\theta) = \mathbb{E}_{\tau \sim \pi_\theta} [R(\tau)]$$

where τ denotes a trajectory of states and actions, and $R(\tau)$ is the cumulative reward of the trajectory. The expectation is over all trajectories possible under policy π_θ .

Policy gradient methods have a number of advantages over value-based methods such as DQN or DDQN:

- They are particularly effective in environments with continuous, high-dimensional action spaces.
- Policy gradient methods can converge to a stable policy due to their gradient-based optimization approach.
- Unlike value-based methods, they can learn stochastic policies with arbitrary probabilities, which are crucial in environments where randomness plays a role in optimal decision making. They are more flexible than ϵ -greedy policies over action values in approaching deterministic policies.

On the other hand, policy gradient methods also have disadvantages, such as:

- The estimates of the gradient can have high variance, leading to inefficient learning and the need for variance reduction techniques.
- They often require a large number of samples to converge, making them inefficient.
- The performance of the policy can be heavily dependent on the initial parameter settings.

A simple policy gradient method is REINFORCE. The REINFORCE method uses the following parameter update method. The update is proportional to the return G_t and inversely proportional to the action probability π .

$$\theta_{t+1} = \theta_t + \alpha G_t \frac{\nabla \pi(A_t | S_t, \theta)}{\pi(A_t | S_t, \theta_t)} \quad (19.2)$$

A complete Monte Carlo based REINFORCE algorithm is shown in Figure 19.4. The basic structure is similar to the tabular MC control introduced in the previous chapter. Complete episodes are generated and the updates are based on the actual return

```

Input: A differentiable policy  $\pi(a|s, \theta)$ ; step size  $\alpha > 0$ 
Initialize policy parameters  $\theta \in \mathbb{R}^d$  arbitrarily
Loop forever (for each episode):
    Generate an episode  $S_0, A_0, R_1, \dots, S_{T-1}, A_{T-1}, R_T$ ,
    Loop for each step of the episode  $t = 0, 1, \dots, T - 1$  :
        
$$G \leftarrow \sum_{k=t+1}^T \gamma^{k-t-1} R_k$$

        
$$\theta \leftarrow \theta + \alpha G \nabla \ln \pi(A_t | S_t, \theta)$$


```

Figure 19.4: REINFORCE: Monte-Carlo Control (episodic) (Source: SB)

G at each step. There is no bootstrapping of estimates using other estimates. The primary difference to tabular MC is in the update step that uses the REINFORCE update formula Equation 19.2.

REINFORCE with Baseline

An extension to the basic REINFORCE method is to use "baselines", values relative to which the return G_t is evaluated. This reduces the variance of the updates but leaves the expected values unchanged, that is, it is unbiased. Additionally, this has been shown to improve the speed of learning.

The main idea is to use the following update that includes a "baseline" return $b(S_t)$ for state S_t :

$$\theta_{t+1} = \theta_t + \alpha(G_t - b(S_t)) \frac{\nabla \pi(A_t | S_t, \theta)}{\pi(A_t | S_t, \theta)}$$

One can choose $b(S_t) = \hat{v}(S_t)$, that is to use the state value function as baseline. This yields the following update:

$$\theta_{t+1} = \theta_t + \alpha(G_t - \hat{v}(S_t)) \frac{\nabla \pi(A_t | S_t, \theta)}{\pi(A_t | S_t, \theta)}$$

A complete implementation is shown in Figure 19.5. Note that the state value function is also a parameterized function, with parameter vector w . The update step must not only update the policy parameters θ but also the value function parameters w . The parameters w are updated using an update step analogous to that of the DQN (Equation 19.1) but for the state value function, rather than the action value function.

```

Input: A policy  $\pi(a|s, \theta)$ ; step size  $\alpha_\theta > 0$ 
Input: A state-value function  $\hat{v}(s, w)$ ; step size  $\alpha_w > 0$ 
Initialize parameters  $\theta \in \mathbb{R}^d$ ,  $w \in \mathbb{R}^d$  arbitrarily
Loop forever (for each episode):
    Generate an episode  $S_0, A_0, R_1, \dots, S_{T-1}, A_{T-1}, R_T$ ,
    Loop for each step of the episode  $t = 0, 1, \dots, T - 1$  :
         $G \leftarrow \sum_{k=t+1}^T \gamma^{k-t-1} R_k$ 
         $\delta \leftarrow G - \hat{v}(S_t, w)$ 
         $w \leftarrow w + \alpha_w \delta \nabla \hat{v}(S_t, w)$ 
         $\theta \leftarrow \theta + \alpha_\theta G \nabla \ln \pi(A_t | S_t, \theta)$ 

```

Figure 19.5: REINFORCE with Baseline (Source: SB)

Actor-Critic Methods

The policy gradient methods in Figures 19.4 and 19.5 are both Monte Carlo methods. Recall that moving from Monte Carlo method to TD methods involved recognizing that $G_{t+1} = R_t + \gamma G_t$ and that the expected values of G_t is the state value of state S_t . Starting with the REINFORCE with baseline update function, the same considerations apply to policy gradient methods.

$$\begin{aligned}
\theta_{t+1} &= \theta_t + \alpha(G_t - \hat{v}(S_t)) \frac{\nabla \pi(A_t | S_t, \theta)}{\pi(A_t | S_t, \theta_t)} \\
&= \theta_t + \alpha(R_{t+1} + \gamma \hat{v}(S_{t+1}, w) - \hat{v}(S_t, w)) \frac{\nabla \pi(A_t | S_t, \theta)}{\pi(A_t | S_t, \theta_t)} \\
&= \theta_t + \alpha \delta_t \frac{\nabla \pi(A_t | S_t, \theta)}{\pi(A_t | S_t, \theta_t)}
\end{aligned}$$

Here, δ_t is the TD error. The resulting "*One-Step Actor-Critic*" method uses bootstrapping to estimate the state value function, that is, it uses estimated values rather than actual returns. Just like TD, SARSA and Q-learning for tabular methods, actor-critic methods can improve on the slow learning of Monte Carlo methods and are also useful for non-episodic, continuous problems. Figure 19.6 shows the complete algorithm for the one-step actor-critic method.

```

Input: A policy  $\pi(a|s, \theta)$ ; step size  $\alpha_\theta > 0$ 
Input: A state-value function  $\hat{v}(s, w)$ ; step size  $\alpha_w > 0$ 
Initialize parameters  $\theta \in \mathbb{R}^d$ ,  $w \in \mathbb{R}^d$  arbitrarily
Loop forever (for each episode):
    Initialize  $S$  (first state of episode);  $I \leftarrow 1$ 
    Loop while  $S$  not terminal (for each time step):
        Sample  $A$  from  $\pi(\cdot|S, \theta)$ 
        Take action  $A$ , observe  $S', R$ 
         $\delta \leftarrow R + \gamma\hat{v}(S', w) - \hat{v}(S, w)$ 
         $w \leftarrow w + \alpha_w \delta \nabla \hat{v}(S_t, w)$ 
         $\theta \leftarrow \theta + \alpha_\theta G \nabla \ln \pi(A_t|S_t, \theta)$ 
         $S \leftarrow S'; I \leftarrow \gamma I$ 

```

Figure 19.6: One-Step Actor-Critic algorithm (Source: SB)

19.5 Additional Information

Stable Baselines

OpenAI Stable Baselines is a collection of RL algorithm implementations. It provides a set of high-quality, efficient, and easy-to-use Python implementations of several state-of-the-art reinforcement learning algorithms. The primary goal of Stable Baselines is to make it simpler for the research community and industry practitioners to replicate, refine, and deploy RL solutions. Stable Baselines has several features that are designed to enhance the usability and performance of RL algorithms:

- *Unified Structure*: Each algorithm adheres to a consistent structure, making it easy to understand, modify, and experiment with different algorithms.
- *Pre-configured Hyperparameters*: It comes with expert-selected hyperparameters that work well out of the box for many problems, reducing the need for extensive tuning.
- *Extensive Documentation and Examples*: Comprehensive documentation and a variety of examples are provided, facilitating quick learning and implementation.

Stable Baselines includes a wide array of RL algorithms, each tailored for different kinds of RL problems. Some of the notable included algorithms are:

- *Proximal Policy Optimization (PPO)*: A policy gradient method that balances the benefits of on-policy and off-policy learning, offering both robustness and stability in performance across a variety of environments.
- *Deep Q-Network (DQN)*: An off-policy algorithm that uses a deep neural network to approximate the Q-value function, suitable for discrete action spaces.

- *Soft Actor-Critic (SAC)*: An actor-critic method that optimizes a stochastic policy and aims for maximizing expected return while also maximizing entropy, making it effective for continuous action spaces.
- *A2C and A3C*: Synchronous (A2C) and Asynchronous (A3C) Advantage Actor-Critic methods that use multiple workers to explore the environment and learn more efficiently.

<https://stable-baselines.readthedocs.io/en/master/>

Gymnasium

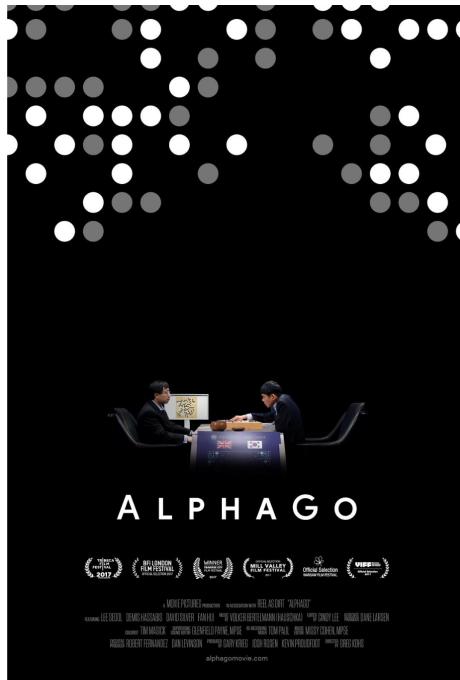
Farama Gymnasium extends the OpenAI Gym framework, providing a suite of RL environments designed for research and education. The Gymnasium environments range from simple toy problems to complex simulations that mimic real-world scenarios. Farama Gymnasium offers several features that make it a useful resource:

- *Wide Range of Environments*: Includes classic control tasks, algorithmic tasks, Atari games, and physical simulations.
- *Standardized APIs*: Maintains consistent APIs across different environments, facilitating easy integration and experimentation with various RL algorithms.
- *Customization and Extensibility*: Allows for customization of environments and easy addition of new ones, enabling researchers to test algorithms on tailor-made scenarios.
- *Community-Driven*: Open-source and community-driven, which encourages contributions and continuous improvement.

The environments in Farama Gymnasium can be categorized into several types, each suited for specific aspects of reinforcement learning:

- *Classic Control*: Simple mechanics and dynamics, such as CartPole, MountainCar, and Pendulum, which are excellent for initial algorithm testing and teaching fundamentals.
- *Atari Games*: Emulated Atari 2600 video games, providing a range of challenges from simple to complex decision-making and control in pixel-based environments.
- *Algorithmic Tasks*: Environments that require the agent to learn underlying algorithms to perform tasks like sorting numbers and simple arithmetic.
- *2D and 3D Robots*: Simulations of robotic tasks including walking, picking, and moving objects, which are more complex and require continuous control strategies.

<https://gymnasium.farama.org/index.html>



<https://www.alphagomovie.com>

Figure 19.7: AlphaGo – The Documentary

AlphaGo

AlphaGo is a significant achievement in the field of artificial intelligence, developed by Google DeepMind. It was designed to play the ancient board game Go, which is known for its deep strategic complexity. AlphaGo's architecture showcases the potential of deep learning and reinforcement learning techniques. AlphaGo combines advanced machine learning techniques, including deep neural networks and Monte Carlo tree search (MCTS). Its design consists of several key components:

- *Policy Networks*: These networks were used to predict the next move during a game. AlphaGo was trained on both human expert games and games it played against itself.
- *Value Networks*: This network predicted the winner of the game from the current position, assisting AlphaGo in evaluating board positions.
- *Monte Carlo Tree Search*: MCTS was utilized to simulate various possible future game scenarios, guiding the policy and value networks to explore the most promising moves further.

The award-winning full-length documentary "AlphaGo" (Figure 19.7) chronicles the journey of the AI program from its initial development through its historic 2016 match

against Lee Sedol, one of the world's top Go players. It provides an in-depth look at the human and technical narratives behind AlphaGo's development. The film highlights several key aspects of RL.

<https://www.alphagomovie.com>

<https://www.youtube.com/watch?v=WXuK6gekU1Y>

The introductory paper on AlphaGo by David Silver and others in the journal Nature should be easy to understand: "Mastering the game of Go without human knowledge". Nature. 550 (7676): 354–359

19.6 Additional Learning Materials

Many well-known and well-published researchers and many companies are actively providing learning materials that can be used to supplement this chapter. They range from introductory materials to full courses on reinforcement learning and are freely available. These researchers and organizations are at the forefront of RL research and the following materials are immensely helpful in understanding this topic.

David Silver Dr. David Silver of University College London is also a lead researcher with Google DeepMind and contributed extensively to the AlphaGo team. He has an excellent introductory course on reinforcement learning with class materials (from 2015) and lectures in a YouTube playlist. Updated courses (2018, 2021) are available on the DeepMind YouTube channel. The 2021 course include topics on deep reinforcement learning.

<https://www.davidsilver.uk/>

<https://www.davidsilver.uk/teaching/>

<https://www.youtube.com/playlist?list=PLqYmG7hTraZDM-OYHWgPebj2MfCFzF0bQ>

https://www.youtube.com/@Google_DeepMind/playlists.

UC Berkeley UC Berkeley hosted a Deep RL Bootcamp in 2017 with slides and lecture videos available online. Additionally, UC Berkeley's course on Deep RL is available online, with lecture slides and videos of past years.

<https://sites.google.com/view/deep-rl-bootcamp/lectures>

<https://rail.eecs.berkeley.edu/deeprlcourse/>

Denny Britz Formerly at the Google AI team, Denny Britz applied RL algorithms to financial markets and trading. He has a interesting blog, and a GitHub repository with resources and algorithm implementations of popular RL algorithms.

<https://dennybritz.com/>

<https://github.com/dennybritz/reinforcement-learning>

Massimiliano Patacchiola Dr. Patacchiola is a postdoc at Cambridge University. He has written a series of excellent blog posts on reinforcement based on the book "Artificial Intelligence – A Modern Approach" by Russell and Norvig. There are lots of illustrations and pointers to implementation and code in multiple languages.

<https://github.com/mpatacchiola/dissecting-reinforcement-learning>

Pascal Poupart Dr. Poupart of the University of Waterloo has made available videos and all course materials for all lectures for a course on reinforcement learning at UWATERLOO.

<https://www.youtube.com/playlist?list=PLdAoL1zKcqTXFJniO3Tqgn6xMBBL07EDc>

<https://cs.uwaterloo.ca/~ppoupart/teaching/cs885-spring18/schedule.html>

Andrew Ng Dr. Ng of Stanford University was the former head of Google Brain and chief scientist at Baidu. He has taught an introductory class on reinforcement learning, as part of a broader course on machine learning.

<https://www.andrewng.org/>

<https://www.youtube.com/watch?v=RtxI449ZjSc>

<https://www.youtube.com/playlist?list=PLA89DCFA6ADACE599>

Andrej Karpathy Andrej Karpathy was a founding member of OpenAI (makers of ChatGPT and Dall-E) and later became the Tesla lead for their Autopilot autonomous driving program. An early blog post by Andrei Karpathy on RL is at the introductory level.

<https://karpathy.ai/>

<https://karpathy.github.io/2016/05/31/rl/>

Lilian Weng Dr. Weng is a lead researchers at OpenAI (makers of ChatGPT and Dall-E). She has written an early blog post on RL and another one on policy gradient algorithms.

<https://lilianweng.github.io/>

<https://lilianweng.github.io/posts/2018-02-19-rl-overview/>

<https://lilianweng.github.io/posts/2018-04-08-policy-gradient/>

OpenAI OpenAI (makers of ChatGPT and Dall-E) post regularly on their blog, on all things deep learning and also reinforcement learning. The blog posts are easy introduction to a variety of analytics topics.

<https://openai.com/blog/openai-baselines-ppo/>

<https://openai.com/blog/evolved-policy-gradients/>

<https://openai.com/blog/evolution-strategies/>

19.7 Review Questions

Introduction

1. Explain the concept of function approximation in the context of reinforcement learning. How does it address the scalability issues faced by tabular methods?
2. How does function approximation help in generalizing from seen to unseen states?
3. Can decision trees be used as function approximators in RL? Discuss their potential advantages and limitations if used.
4. Describe how function approximation can be applied to state values, action values, and policies. Provide the mathematical representation used for each.
5. How does function approximation contribute to the flexibility and efficiency of reinforcement learning models?
6. Provide an example scenario in reinforcement learning where generalization from seen to unseen states would be crucial.
7. Discuss the issues of stability and convergence in function approximation methods, especially when combined with off-policy learning.

8. What measures can be taken to prevent overfitting in function approximation models, particularly those using deep neural networks?
9. Given the advantages and challenges of function approximation, in what types of reinforcement learning problems would you recommend its use?
10. Imagine you are designing a function approximation model for an RL problem in a financial trading environment. What factors would you consider in choosing the type of function approximator?

Value-Based Methods and Stochastic Gradient Descent

11. What is the formula for the value error (VE) in the context of action values and how is it computed?
12. Derive the gradient of the MSE loss function used in the context of function approximation for reinforcement learning.
13. Describe the parameter update rule in stochastic gradient methods for function approximation. What does each term in the update equation represent?
14. What is bootstrapping in the context of reinforcement learning? How is it implemented in SGD updates?
15. Describe the concept of experience replay and its significance in stabilizing the SGD updates in reinforcement learning.
16. Discuss the role of the target network in the Double Q Network (DQN) algorithm. How does it contribute to the stability of the learning process?
17. Compare the update steps in tabular SARSA and semi-gradient SARSA using function approximation. What is the key difference?
18. Explain how the stochastic gradient SARSA algorithm is adapted to utilize a replay buffer and target network in the context of the DQN algorithm.
19. What are the components of the "deadly triad" in reinforcement learning? Describe how each component contributes to instability and divergence.
20. Provide examples of how modern reinforcement learning algorithms address the challenges posed by the deadly triad.
21. Explain the impact of periodic updates from the main network to the target network. How does this timing affect the algorithm's performance?
22. In the context of function approximation, how is the learning process affected when using non-linear function approximators like neural networks compared to linear approximators?

Policy Gradient Methods

23. Describe how policy gradient methods optimize the policy parameters directly. What is the significance of this approach in environments with continuous action spaces?
24. Explain the typical representation of a policy in policy gradient methods and how it relates to the probability of selecting actions.
25. Define the objective function $J(\theta)$ used in policy gradient methods. What does this function represent?

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26. List and describe the main advantages of using policy gradient methods over value-based methods in reinforcement learning.
27. Explain the principle behind the REINFORCE algorithm. How does it update the policy parameters?
28. Describe the update rule of the REINFORCE method. How does the inclusion of the logarithm of the policy's probability function influence the update?
29. Explain why the REINFORCE algorithm updates parameters only at the end of each episode. What are the limitations of this approach?
30. What is the purpose of using a baseline in the REINFORCE algorithm? How does it affect the variance of the updates?
31. Explain the update formula used in REINFORCE with baseline. How does the inclusion of the baseline value $b(S_t)$ change the update mechanism?
32. Compare the REINFORCE algorithm to Actor-Critic methods. How do Actor-Critic methods improve upon the basic policy gradient approach?
33. How does the One-Step Actor-Critic algorithm use the current and next state values to update the policy and value function parameters?
34. Explain how the Actor-Critic method combines the benefits of policy gradient and value function approximation methods. What are the specific roles of the "actor" and the "critic"?

Chapter 20

Managing Machine Learning Operations (MLOps)

Sources and Further Reading

The material in this chapter is based on the following sources.

Treveil, M. and the Dataiku Team (2020) *Introducing MLOps*, O'Reilly Media, Sebastopol, CA (T)

The book by Mark Treveil and others is a good, short introduction to the principles and ideas of machine learning operations. It provides guidance on the process, the management, and the governance of MLOps. Given its focus on high-level introduction, it provides few technical details, but this also means that this book will likely stay relevant longer.

Gift, N. and Deza, Al. (2021) *Practical MLOps*, O'Reilly Media, Sebastopol, CA (GD)

The book by Noah Gift and Alfredo Deza provides a more "hands-on" introduction to machine learning operations. While also touching on the principles and processes, it goes in-depth and offers specific technical illustrations of good MLOps practices. The book uses both on-premises technology and cloud-based technologies, but focuses most heavily on MLOps on the AWS cloud.

Resources

Complete implementations of all examples in this chapter are available on the following GitHub repo:

<https://github.com/jevermann/busy4720-mlops>

The project can be cloned from this URL:

<https://github.com/jevermann/busy4720-mlops.git>

20.1 Introduction

Many introductions to machine learning and predictive models in particular focus on the different types of statistical models, their properties and relative advantages, and challenges and best-practices for feature engineering and model training. Using the R or Python environment, the focus is often on the business analytics team that identifies features and creates the model. A frequent implicit assumption is that this happens in isolation, on relatively small data sets, and on desktop computers. Often, such an introduction fails to examine not only infrastructure challenges but, more importantly, also neglects to examine how the created models are managed and used productively. The latter is arguably most important from a business perspective.

As an example, consider a dynamic pricing model that predicts what customers are willing to spend on a particular product based on their previous purchase history and other features. It is relatively easy to collect a small data set and train a few different models. However, ultimately, this model must be integrated with the web-based store front of the organization and will be used to show different prices to different customers. It is a long way from a small model on the desktop to a model that is in production use within an organization's larger set of software applications. Besides problems of efficiently moving models to production, that is, "*deploying*" them, models in production also pose certain risks to an organization and these risks must be managed.

Purpose

Machine Learning Operations, or MLOps, therefore has three main purposes. First, it seeks to improve operational efficiency in model management and deployment. It does this through formalized and automated processes to achieve reliability and repeatability in deployment.

The second purpose is to manage and mitigate risk. Different types of risks arise, such as availability of service. What happens when the dynamic pricing model in the example becomes unavailable due to a computer outage or a software misconfiguration? Will the organization lose money? Will it lose customers? Another risk stems from the model quality and the impacts of model predictions. For example, how large are the prediction errors of the dynamic pricing model and how much potential revenue could

the organization lose by overpricing items and causing the customers not to purchase something? How much potential revenue remains unrealized because items are offered at a lower price than a customer would be willing to pay? A third type of risk arises from prediction fairness. For example if the pricing model includes features such as gender so that different prices are offered to men and women, would that be fair (or even legally allowed)? If this were to become widely known to customers, would customers defect from the business? Finally, there is the risk of skill loss. What happens when the business analysts who developed the model leave the organization? Does the organization have sufficient documentation for auditability and risk management? Can the model be recreated by a new team? How easily can a replacement team member become familiar with a model and its history, rationale, and current uses?

The third purpose of MLOps is to establish accountability, auditability, and traceability. Accountability is concerned with decisions made about models, their acceptance, and their deployment into production. It requires formalized testing and acceptance procedures for the model as it moves from data extraction to deployment. This begins with questions whether there is legal or ethical clearance to use a particular data set, to the decision as to when a new model replaces an existing model in a production environment, e.g. in the web-based store front of the dynamic pricing example. Auditability and traceability have two aspects. First, they refer to internal processes and procedures and concern the ability to answer questions about the origins of the training data, of the model parameters, the production environment, and related decisions, for example about feature inclusion or risk assessment. A second aspect is the auditability and traceability of model output and/or decisions made based on the model output. In the dynamic pricing model example, the organization must maintain a record of why a certain price was offered to a particular customer, what the relevant features were that led to the decision, and be able to trace these features back through the model all the way to the training data.

Challenges

As business analytics, machine learning, and predictive models have become popular, organizations face a number of challenges. First, the popularity and the competitive necessity for organizations to engage in this area often leads to a proliferation of separate business analytics teams in different organizational units, that produce a multitude of models, from a wide variety of data, for a large number of potential use cases. In many organizations, machine learning is not well organized and managed, leading to conflicting goals, unanticipated model interactions, lack of traceability and auditability, lack of compliance, lack of knowledge of model performance, lack of accountability and, in the worst case, customer-facing problems where the organization's customers that are affected by a predictive model's decisions become aware of its poor quality.

A second challenge is the fact that data is constantly changing. After a model has been trained, the organization continues to receive or collect data that can and should be used for model training. If nothing else, additional training data can reduce the prediction error of the model. However, the main challenge is that data may change over time. For example, the business may attract customers with different characteristics and needs, or

the industry competitive position of a business may change so that customers have different preferences, or the macro-economic conditions may change to make customers more risk averse or less willing to spend money. These challenges can lead to models that perform poorly, and require a systematic approach to model performance monitoring, model retraining, and tracking the relative performance of different models and different model versions.

As a third challenge, the needs of the business or organization can change over time, making predictive models irrelevant or ill-suited for their purpose. Organizations may shift their strategies, enter or exit specific markets, adapt their marketing strategies and tactics, change their business processes and operations and many other aspects. Such changes will affect the usefulness of predictive models that are deployed in the organization and may require replacing or retraining models, perhaps with new input features, different targets, and with trained with loss functions more appropriate to the changed business goals and objectives.

A fourth challenge is the mixed composition of business analytics teams. Such teams are often comprised of business professionals as subject matter experts, data scientists that focus on data quality, data provenance, and model development, software engineers that must take a trained model and embed it into an organization's software applications, for example, into their web-based sales tool or their mobile app. Finally, IT staff participate on teams to provision appropriate infrastructure and manage IT related risk. The problem with such interdisciplinary teams is often that people may be unaware of the roles or the importance of others on the team, they may be focused on optimizing their particular aspect of model development to the detriment of the overall effort, they may use different terminology (for example, what precisely does the term "model" mean to each of them?), and they may not have developed effective management processes or means of communication.

Finally, data scientists tend to have little expertise in software engineering and software deployment. Traditionally, this role has focused on the specific details of the statistical models without any reference to established practices that makes software products reliable, robust, scalable, easily maintainable, auditable, fault and failure tolerant, safe, or a variety of other desirable properties. However, when machine learning models or predictive or prescriptive models and algorithms are to be deployed within an organization or even facing an organization's customers, the models become part of an organization's software and therefore such software properties become very important, possibly more important than the predictive accuracy of a machine learning model.

Principles

MLOps is a set of practices based on a few main principles. The first principle is that of reliability and reproducibility of operations. This requires standardized and structured processes that are made explicit, either in the form of documentation or, if they are automated, in the form of programming code. Such a process typically includes the steps beginning with data extraction from operational systems, data preprocessing and cleaning, feature extraction and feature engineering, training and evaluation of multiple

models, hyper-parameter search, automated model testing, automated software building, and ending with automated deployment. Each process step must be documented and the results archived for auditability purposes.

It is clear that describing the process with executable code, e.g. in the form of Python scripts or using a variety of open-source or cloud-based products, is preferable to simply writing out a procedure manual. This preference for code illustrates the second main principle of MLOps, that of robust automation. Together, the first two principles ensure that every model in production can be easily re-created in identical form when needed. More importantly, for auditability purposes, the principles ensure that the model and data provenance are captured and documented, that decisions and actions as to which model to move into production are documented, and that the organization knows which models are currently in production and what their properties (e.g. error rate, cost, resource consumption, risk assessment results, etc.) are.

A frequently used term related to robust automation is that of "infrastructure as code". Modern development, testing, and production environments are often provisioned automatically. That is, computer servers, software, databases, file storage are provided automatically as needed. To enable this, the requirements for a particular model and how to provide or create them must be described. To take one example, building and deploying a predictive model as a micro-service requires instructions on how to build the software that serves the model predictions (e.g. a Makefile¹), instructions on how to build the container that the software will run in (e.g. a Dockerfile²), instructions on what type of machine to run the container on, etc. These instructions are themselves computer code and they must be managed, versioned and tested as computer code.

The third principle is that all versions of both the data and the models must be maintained and managed. Models in this context mean more than just the final parameter values but includes evaluation results, hyper-parameters, training and testing history, model explainability results, model rationale, versions of all software packages that were used, and any other documentation. A variety of open-source and commercial on-premises and cloud-based tools are available for this, ranging from software code versioning tools like git and GitHub to model registries on Amazon Web Services or Microsoft Azure. Versioning of the training data is more problematic, as training data sets can be very large. It is often simply not feasible to keep copies of different versions and more intelligent approaches are being developed. Managing and versioning data and models are useful both for automating model deployment and for ensuring auditability and through this, for ensuring compliance with internal or external requirements. For example, model versioning and management allows an organization to say

¹A Makefile contains computer readable instructions in a standardized format that allows the "make" software tool to automatically build and test a software application. For more details, see this page: [https://en.wikipedia.org/wiki/Make_\(software\)](https://en.wikipedia.org/wiki/Make_(software)).

²Containers are "light-weight" virtual machines with their own operating system, databases, and other software installed in them. Docker is one particular container technology. A Dockerfile defines how to build a docker container, that is, what operating system to use, which software packages to install and how to configure the software in the container for use. For details on containerization see this page: [https://en.wikipedia.org/wiki/Containerization_\(computing\)](https://en.wikipedia.org/wiki/Containerization_(computing)) and for detail on Docker, see this page: [https://en.wikipedia.org/wiki/Docker_\(software\)](https://en.wikipedia.org/wiki/Docker_(software)).

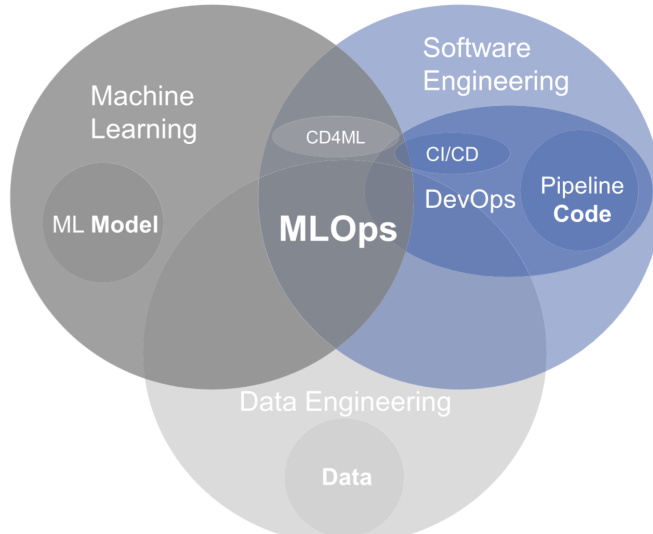
precisely what features and input data was used to train a model, and what features and their relative importance are being used to make decisions.

The fourth principle of MLOps is continuous delivery to production. The idea is that rather than treating model development and deployment a single or one-off project, models should be continuously improved and the improved models should be continuously moved into production. That is, as new training data becomes available, model training continues or the existing model can be fine-tuned with the new data. Of course, continuous delivery and integration does not mean to do this whenever a single new training observation becomes available, but to have an established frequency, depending on model size and required training time, of doing continuous delivery daily, weekly, monthly, quarterly or annually. Importantly, training or fine-tuning the model is just the initial step that kicks off the delivery process. Activities like testing, risk assessment, documentation, performance comparison, software integration, etc. need to follow for each newly created model.

Finally, to ensure traceability, auditability, risk management and identify when models should be replaced, retrained or fine-tuned, continuous monitoring of a model in production is required. This means that all inputs and all outputs to the model must be logged and analyzed. In the dynamic pricing model example, this means that whenever a customer is offered a price for an item that is determined by the predictive model, all relevant input features and the model prediction must be logged, as well as the actual purchase decision or other resulting action by the customer. It is obvious that such log data can grow quickly in size and appropriate infrastructure needs to be in place to manage the data. More importantly, procedures and processes need to be in place to actively monitor and analyze this data to detect changes in customer features over time or changes in model accuracy over time.

Relationship to Other Disciplines

Machine learning operations (MLOps) is at the intersection of a number of related disciplines, as shown in Figure 20.1. It overlaps with machine learning where model building, training, evaluation, etc. are located ("ML Model" in Figure 20.1). Specifically, MLOps focuses on that part of machine learning that is sometimes called "CD4ML" (continuous delivery for machine learning), that is, the software engineering principles of continuous delivery as applied to machine learning models. MLOps overlaps with software engineering, which focuses on continuous integration and continuous delivery ("CI/CD" in Figure 20.1, "DevOps" (unified or integrated development and operations), and automation of software delivery pipelines. Finally, MLOps overlaps with data engineering, which focuses on collecting, managing, and providing data for organizational purposes, including but not only to machine learning applications. Other uses of data in business analytics may be databases for descriptive analytics or visualizations.



Source: Kreuzberger, D., Kühl, N., & Hirschl, S. (2023). Machine learning operations (mlops): Overview, definition, and architecture. *IEEE access*, 11, 31866-31879.

Figure 20.1: MLOps – Relationship to other disciplines

20.2 MLOps Lifecycle Overview

The MLOps lifecycle is a combination of the machine learning model lifecycle, shown in Figure 20.2, and the software development "DevOps" lifecycle in Figure 20.3. The resulting combination is shown in Figure 20.4.

The model development lifecycle in Figure 20.2 contains activities to manage the data and to develop a machine learning or prediction model. Data collection can include activities such as ETL ("extraction, transformation, loading") data from source systems, acquisition from external data brokers or vendors, or scraping data from web sites or news sites, or any number of other ways. The data must be curated. For example, its provenance must be established and recorded, its quality must be assessed, legal or licensing issues must be resolved, etc. The data can then be transformed. This includes both pre-processing for data cleaning as well as engineering new features or combining multiple data sets. Data validation then ensures that the data is of high quality, relevant to the problem, fit for purpose and can legally be used.

The model part of the model development lifecycle begins with data exploration, that is, gaining an understanding of the various features, their interactions, summary statistics, etc. This leads to defining and training one or more models on the data set. This step can also include model search (for example, varying the architecture of a neural network model) or hyper-parameter search (for example, finding the optimal depth of a decision tree). The models are then evaluated and compared on suitable metrics, using hold-out samples or cross-validation. Finally, the evaluation results can be used

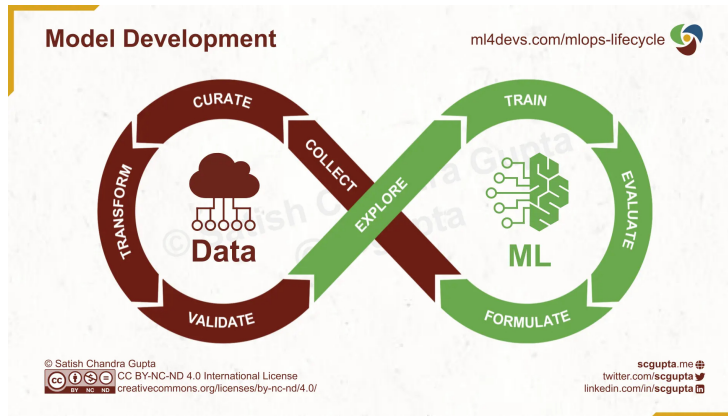


Figure 20.2: Model Development Lifecycle

to inform additional data collection or additional features to be used in a better model leading to another iteration of lifecycle.

DevOps (that is, integrated development and operations) is an approach to software development that focuses on continuous integration and continuous deployment. In early software development, a software application was created by developers, typically as a one-off project, and then turned over to the IT operations department to put it into production. Having separate teams for development and operations leads to problems and frictions. For example, the development team might not be concerned with how much resources their application will require or what operational or security risks an application poses. This puts significant pressure on the operations team. The DevOps approach to software development focuses on an integrated process and integrated teams. Rather than developing software in large one-off projects, DevOps encourages continuous improvement.

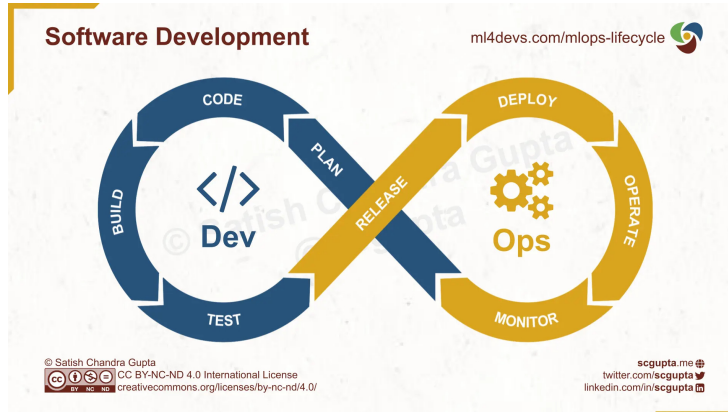


Figure 20.3: Software Development Lifecycle

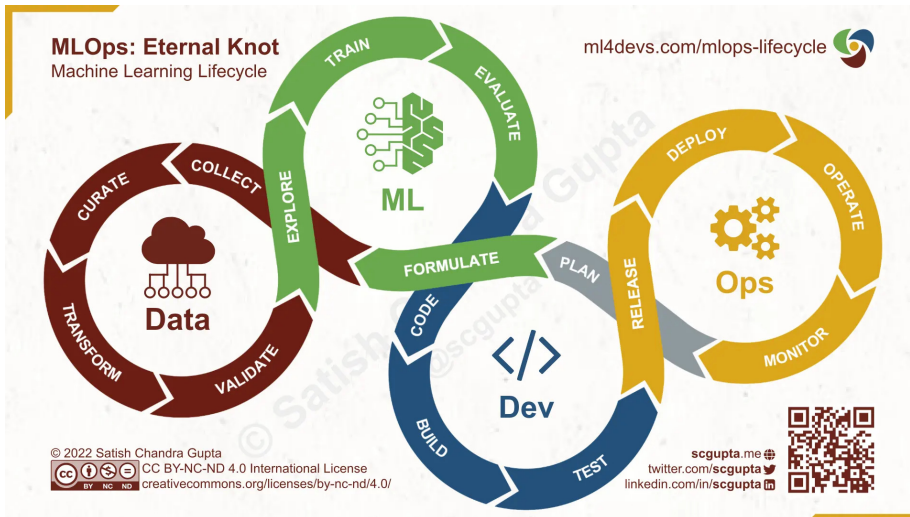


Figure 20.4: MLOps Lifecycle

The lifecycle in Figure 20.3 begins with planning the software application, defining what the application is required to do, its users, its business objective, etc. Software developers then produce programming code. The build phase integrates the separate code pieces from all developers into a complete application, and from this builds the actual software application in the development environment. The test phase moves the software application to a testing environment. It conducts functional tests to ensure the application works correctly, but also performs security tests and performance tests to ensure it poses little risk and consumes reasonable resources. An application is then formally released into production. Resources such as hardware, databases, file storage space, network connections, etc. are provisioned, forming the production environment. The application is then deployed into its production environment, integrated with other software applications and operated. Continuous monitoring ensures the software applications continues to work, to work correctly, and to consume only the anticipated resources. The analysis of monitoring data will lead into a new iteration of the DevOps lifecycle where the next version of the application is planned, developed, and operated.

The combined lifecycle model in Figure 20.4 recognizes the fact that machine learning models are just one part of a complex software application. Consider the example of the dynamic pricing web-store application. The actual prediction model, while central and important, is but a small part of it that must integrated into the web-store application. When the available training data changes (in the "collect" phase), all phases of the lifecycle may need to be performed again to create a new model, develop or adapt the software application and move it to operations. Given the number of steps involved and their complexity, it becomes clear that managing this lifecycle efficiently, and managing it at scale for multiple models, requires formalized and automated processes.

20.3 MLOps Roles and Requirements

Many people participate in different roles in MLOps processes. They participate in or take responsibility in different phases of the overall MLOps lifecycle, as shown in Figure 20.5. This figure shows a simplified representation of the MLOps lifecycle in just 4 phases that are further explained in Section 20.5 below. Each role in the MLOps lifecycle also has certain requirements in order to perform the role efficiently.

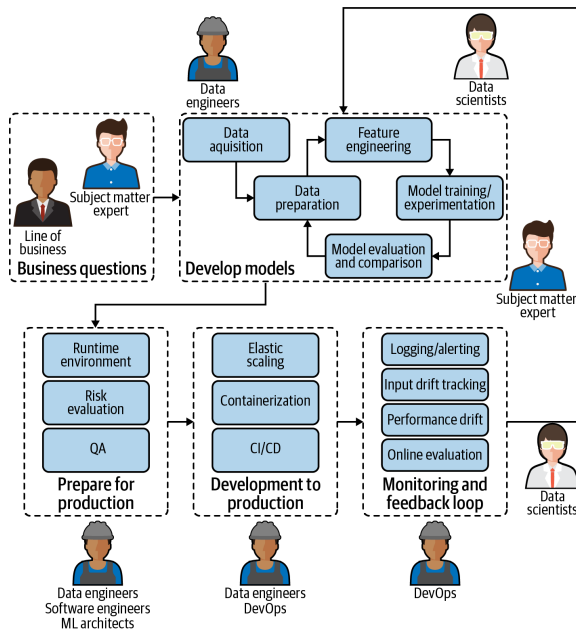
Subject matter experts are business stakeholders that provide the business problems, questions, or goals for a machine learning project. They also define how the success of a project will be measured in business terms by defining the relevant KPIs (key performance indicators). In a dynamic pricing example, a business KPI might be to improve the acceptance rate of customized product offers by 5 percent. These KPIs inform the choice of model and of model evaluation metrics. Once the model is in production, subject matter experts evaluate the performance against the business needs, that is, the KPIs. This evaluation cannot be done by the MLOps team, because it concerns the business goals, not the predictive performance of a model: The model may perform well in predicting the price customers are willing to pay, but customers for other reasons choose not to accept product offers.

Subject matter experts require understandability and interpretability of models, if they are to assess them in business terms. Moreover, they require a responsive feedback mechanism so that their evaluations and assessments can inform the next iteration of the MLOps lifecycle.

Data scientists use available data sets to develop, train, and evaluate different machine learning models, based on the goals defined by the subject matter experts. Their tasks also include related activities such as data preprocessing and feature engineering that are necessary for training and evaluation.

Data Engineers work with data scientists to acquire or collect, manage, clean, maintain, and provide data. They build the data infrastructure necessary for data scientists to efficiently access the data during model training and evaluation. This includes providing file storage space or databases or data warehouses. Data engineers are also involved in data acquisition from internal or external sources, in evaluating and maintaining data quality and in maintaining data provenance.

To perform their tasks efficiently, data scientists and data engineers require automated model packaging and delivery tool. That is, once a model has been developed and found to be have acceptable performance, packaging the model (that is, the trained parameter values, hyper-parameter values, model architecture, training data provenance and description, required software packages, etc.) so that it can be moved to production deployment should be automatic. Data scientists and data engineers also require automatic testing for models. That is, once a model has been trained, it should automatically be subjected to a variety of tests, from simple predictive performance testing



Source: Treveil et al. (2020), Figure 1-3

Figure 20.5: Roles in the MLOps lifecycle

on a range of metrics, through sensitivity testing for a range of normal and abnormal inputs, to interpretability testing.

Data scientists and data engineers require visibility into model performance as the model moves across the development, staging or preparation, and deployment or production phases. That is, they require access to raw performance data but better still, access to online visual dashboards for each model or a set of models. They also require visibility into the data pipeline that is used for each model.

Software Engineers integrate trained and operationalized models into the software applications of an organization. This can take different forms depending on the nature of the software application. For example, a trained model could be integrated into a mobile application, or it could be accessed as a separate service from the organization's web-based store front. They are supported and advised by data engineers and ML engineers or architects in identifying the most efficient way to deploy and deliver the model within the organization's software applications.

Software engineers require programming code versioning systems (such as git or GitHub) to collaborate and track the changes that each member of a team makes to the application's code. They also require automatic testing tools for the software application so that, as they make changes to an application's programming code, their changes can be automatically tested locally for correctness and functionality (unit tests) and in the

large for compatibility with other changes made by the team (integration tests).

DevOps Engineers are responsible for building the system developed by the software engineers and testing them for security, performance, resource use, failure and fault tolerance, and for availability. DevOps engineers perform the CI/CD processes for continuous integration and continuous delivery of software applications through programming code integration, software building, software testing, to software deployment.

DevOps engineers require seamless and automatic deployment pipelines, from software code integration, to final testing, infrastructure provisioning (for example, virtual machines, servers, containers, etc.) to deployment into production. In particular, the MLOps lifecycle should integrate at this point with DevOps lifecycle that may already exists in the IT department of an organization and leverage existing processes and tools.

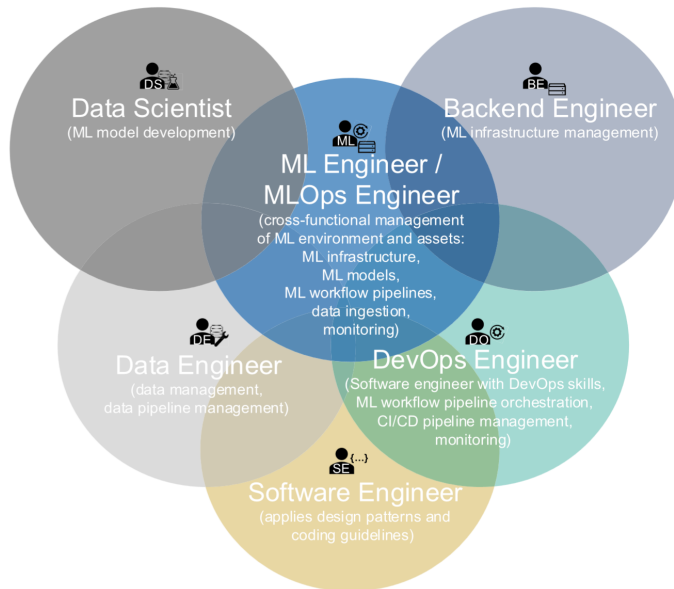
Model risk managers and model auditors are responsible for assessing all types of model risks and ensuring compliance with internal or external regulations and requirements. They assess the models that are developed by data scientists before they move into production. They also monitor the performance of models that are in production to identify relevant changes to the model risk.

Model risk managers require automated reporting on all models (past and present), including data provenance and all model test results.

ML Engineers and ML Architects provide advice on how best to deploy a set of models, the infrastructure required, and the implications of the types of model deployment to software applications. They work closely with data scientists and data engineers who provide knowledge of the model and its data requirements. They make large-scale architectural decisions, such as whether to deploy the model in cloud-based environments like Amazon Web Services or Microsoft Azure, or on-premises, for example using a Kubernetes cluster, or in a web browser or mobile application. They determine scalability and infrastructure requirements.

ML engineers require the ability to easily assess and to quickly adjust infrastructure capacities. For example, if an ML engineer realizes that a ML prediction service performs slowly, the engineer must be able to quickly allocate more servers, deploy additional copies of the ML model, and make them available to the organization's software applications that use that prediction model.

Backend Engineers provide and manage the computational infrastructure. They create and maintain on-premises or cloud-based computer clusters and distributed storage systems. They design and manage fast and efficient network connections between all components, provide backup and restore functionality, and are responsible for high availability of all infrastructure components. Backend engineers are not included in Figure 20.5 but are shown in Figure 20.6 which is taken from a different source.



Source: Kreuzberger, D., Kühl, N., & Hirschl, S. (2023). Machine learning operations (mlops): Overview, definition, and architecture. IEEE access, 11, 31866-31879.

Figure 20.6: Overlapping MLOps Roles

20.4 MLOps Tooling

Because of its heavy focus on automation, MLOps requires a variety of software tools to support it. This section describes important types of software tool and provides references to some popular examples.

Source Code Management (SCM), Code Versioning and Code Repositories are tools that are used by software engineers to collaborate on the creation of a large software application. These tools keep track of all changes and ensure that developers do not overwrite each other's work. The most popular cloud-based tool GitHub³ is based on the open-source git system⁴, but many others exist, both open-source and commercial, cloud-based or on-premises.

CI/CD tools are used to integrate software application programming code, automatically build the software system, test it in a test environment, and then deploy it into a production environments. Popular tools of this type are the open-source Jenkins system⁵ and GitHub actions⁶ on the popular GitHub cloud-based platform.

³<https://github.com>

⁴<https://git-scm.com/>

⁵<https://www.jenkins.io/>

⁶<https://github.com/features/actions>

Workflow Orchestration tools define the usage of data, model, software and configuration artifacts through the development, test, and deployment cycle. They coordinate data extraction, model training, model inference/prediction, and model or software deployment. A popular open-source on-premises system is Apache Airflow⁷, while AWS SageMaker⁸ and Azure Pipelines⁹ are popular cloud-based tools.

Feature Stores offer centralized storage and management of data and any engineered features for ML models. They track feature updates and changes, assess and monitor data quality, and quickly provide data at training or at prediction time. Popular tools are the open source system Feast¹⁰, the AWS SageMaker Feature Store¹¹ or Tecton¹².

Model Training tools support the definition, training, evaluation and comparison of ML models. They are open-source tools like R, Scikit-Learn, TensorFlow, or Spark, or integrated cloud-based offerings such as AWS SageMaker¹³ or Azure Machine Learning¹⁴.

Model Registries provide storage for trained models and their meta-data. This includes the model itself, evaluation results, hyper-parameters, training history, data provenance, documentation, and many other artifacts. These tools provide centralized storage, secure access, version management, and other features. They allow model versions to be tracked and to be quickly deployed in the MLOps lifecycle. A popular open-source model registry is MLFlow¹⁵ while the major cloud platforms also provide model registries, for example the AWS SageMaker Model Registry¹⁶ or the Azure ML Model Registry¹⁷.

ML Metadata Stores store meta data about MLOps pipeline executions, model training, model lineage, etc. For example, they track when a model was trained, evaluated, packaged for production, moved into deployment, etc. These tools support the auditability and traceability of models and their deployment, in turn supporting risk management and compliance assurance.

Model Serving tools execute the model in its production environment and provide access to model predictions or model explanations for the organization's software ap-

⁷<https://airflow.apache.org/>

⁸<https://aws.amazon.com/sagemaker/>

⁹<https://azure.microsoft.com/en-us/products/devops/pipelines>

¹⁰<https://feast.dev/>

¹¹<https://aws.amazon.com/sagemaker/feature-store/>

¹²<https://www.tecton.ai/>

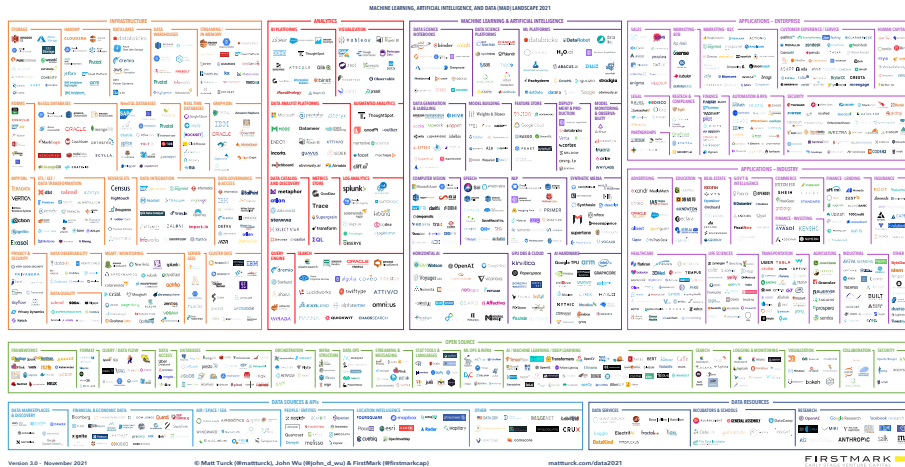
¹³<https://aws.amazon.com/sagemaker/>

¹⁴<https://azure.microsoft.com/en-ca/products/machine-learning>

¹⁵<https://mlflow.org/>

¹⁶<https://docs.aws.amazon.com/sagemaker/latest/dg/model-registry.html>

¹⁷<https://learn.microsoft.com/en-us/azure/machine-learning/how-to-manage-models>



Source: Turck, Matt. *Red Hot – The 2021 Machine Learning, AI and Data (MAD) Landscape*. September 28, 2021.
<https://mattturck.com/data2021/> (last accessed July 22, 2024)

Figure 20.7: Commercial Offerings in the ML Landscape

plications. Popular tools are WSGI¹⁸ servers such as Flask¹⁹, that are often deployed in containers, TensorFlow Serving²⁰, TensorFlow Lite²¹ or TensorFlowJS²² for TensorFlow models, or AWS SageMaker Endpoints²³ for the AWS cloud platform.

Model Monitoring tools can monitor both the computational performance as well as the prediction inputs and outputs of deployed models. Cloud-based examples are the AWS SageMaker model monitor²⁴ on the AWS platform or the Azure ML model monitor²⁵.

As MLOps is a relatively young discipline, tool support, both open-source, commercial, and cloud-based, is still evolving rapidly. Figure 20.7 shows a (partial) overview of companies that offer products and services in this space. Because of the rapid evolution of offerings in MLOps it is simply not possible to provide a detailed description or even a meaningful introduction to these software tools that would remain relevant for more than a few months.

¹⁸ https://en.wikipedia.org/wiki/Web_Server_Gateway_Interface

¹⁹ <https://flask.palletsprojects.com/en/3.0.x/>

²⁰ <https://www.tensorflow.org/tfx/guide/serving>

²¹ <https://www.tensorflow.org/lite>

²²<https://www.tensorflow.org/js>

<https://docs.aws.amazon.com/sagemaker/latest/dg/realtime-endpoints.html>

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²⁴ <https://docs.aws.amazon.com/sagemaker/latest/dg/model-monitor.html>

25 <https://learn.microsoft.com/en-us/azure/machine-learning/>

<https://learn.microsoft.com/en-us/learn/modules/concept-model-monitoring/>

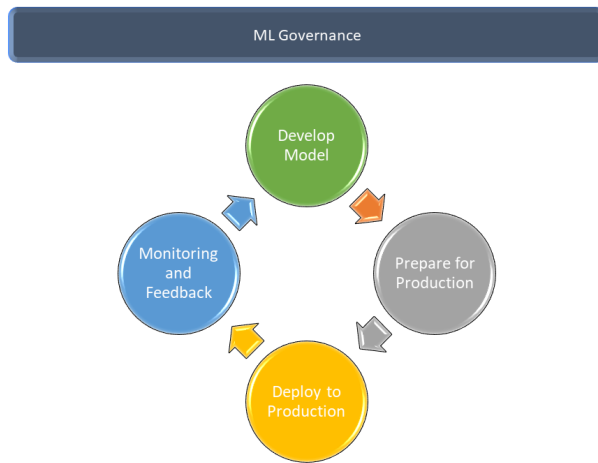


Figure 20.8: Simplified MLOps Lifecycle and ML Governance

20.5 MLOps Lifecycle Phases

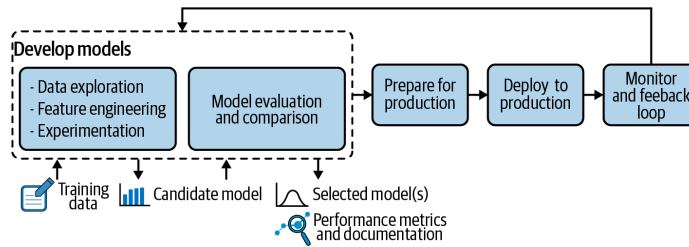
Figure 20.8 shows a simplified MLOps lifecycle together with ML governance. While the MLOps lifecycle concerns operations and management, ML governance refers to the oversight and risk management of the MLOps lifecycle and its processes, participants, and tools. The lifecycle in Figure 20.8 is an abstracted version of the lifecycle underlying Figure 20.5 above. This section provides brief additional comments on each phase in this simplified lifecycle.

20.5.1 Develop Models

The first phase of the simplified lifecycle, develop models, is highlighted in Figure 20.9. It comprises data and feature engineering, as well as model building, training, evaluation, and comparison. The inputs are training data and other models for comparison (e.g. alternative model types, alternative architecture, the current production model, etc.). The output of this phase are a selected model and its performance metrics and associated documentation.

From the data management perspective, important questions to ask and answer in this phase are about data permission or licenses, access requirements, and legal or regulatory obligations or constraints. These are questions that help assess the risk and to assure compliance. Specific example questions are:

- What data are available? What is the quality of that data?
- Can the data legally be used for this purpose? What are the terms of use of the data? What licenses are available or in place for using the data? What is the cost of any required licenses?



Source: Treveil et al. (2020), Figure 4-1

Figure 20.9: Model development in the MLOps lifecycle

- How can the data be accessed? Is it available via an external service or does it exist on-premises? What is the technical access mechanism?
- What features can be created by combining multiple data sets?
- Must the data be redacted or anonymized? Does it include personally identifiable data, such as names or addresses? Does it include sensitive data, such as medical or financial information?
- Are there features that cannot be used legally (age, gender, race, etc.) for some purposes? For example, while age and gender could be used in a dynamic pricing application, they are not legally useable in many jurisdictions for evaluating job applications or making hiring decisions.
- Is the data representative of minority classes/populations? When the data set is biased towards a large majority class or only covers part of the target population, a predictive model built from it will not perform well for minority classes. At best, this increases the risk due to poor predictions, at worst, this prohibits the use of the model in some jurisdictions as it may not exhibit fairness towards all classes or sub-populations.

Tool support for data management in this phase includes ETL pipelines to extract data from source systems, transform it, and load it into a central data or feature store. It also includes centralized data storage, feature storage, and feature engineering tools.

From the model management perspective, important questions also address bias and fairness of model predictions or outcomes. Again, this is to assess and mitigate model risk and to ensure compliance with regulatory and legal obligations. Specific questions to ask include the following:

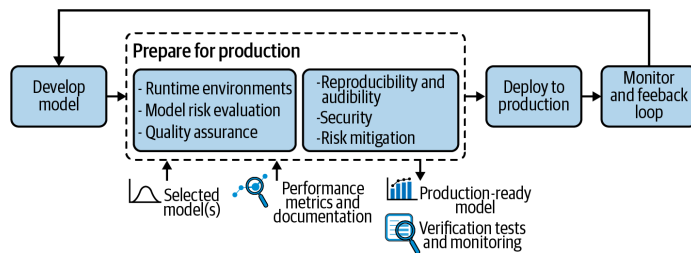
- What are appropriate evaluation metrics? How do these metrics relate to the business KPIs and the goals of the model?
- Is the model performance acceptable globally and for different sub-populations? How are different performance metrics, such as precision and recall, combined into an overall model performance assessment?

- Does the model need to be interpretable or explainable? What type of explanations, for example global or local, should be provided to the model user?
- Are the model outcomes fair to all possible users? How is fairness defined? Are there conflicting definitions or requirements for fairness and how are they combined and reconciled?

Tool support for model definition and management include model registries and repositories that can keep track of and store the models themselves (trained parameters), but also the hyper-parameters, random seeds, software versions, train and test histories and results, associated documentation, etc. Feature stores are used at this stage to provide the data for model training and container makefiles and container registries are often used to provide fixed and reproducible training environments, including all required software packages in their specific versions.

20.5.2 Prepare for Production

Preparing a model for production, highlighted in Figure 20.10, includes selection of the runtime environment and deployment mode (e.g. as a micro-service, as a model embedded in a mobile app, etc.). This decision is made by ML engineers in collaboration with data engineers and data scientists. Preparing for production involves risk assessment of the model and quality assurance of the model by model risk managers and model auditors. Risk mitigation measures are put into place (e.g. an automatic fail-safe when out-of-scope inputs are detected), the model security is tested and improved if necessary, and reproducibility and auditability of the model are confirmed.



Source: Treveil et al. (2020), Figure 5-1

Figure 20.10: Prepare for Production in the MLOps Lifecycle

The input to this phase are the selected model together with its performance metrics and associated documentation. The output is a production ready model with the results of the verification tests performed during this phase.

Technical questions to ask and answer and decisions about the runtime environment to be made during this phase include the following:

- What is the runtime environment and deployment mode? Example options are WSGI services in containers deployed on clusters (on-premises or cloud-based),

TensorFlow Serving deployment (again, either on-premises or cloud-based), so-called "edge-devices" such as mobile phones, embedded systems or IoT devices (internet-of-things), or in a web-based application as a JavaScript model.

- Does the model need to be adapted for production? For example, quantization can increase the performance of model predictions significantly by replacing large floating point numbers with smaller ones. Often, models are trained with 32 bit floating point numbers, but then deployed with 8 bit numbers. The loss in precision may be negligible while the performance benefits can be significant. Another example of model adaptation is additional pruning of decision trees. Again, careful pruning may have a negligible impact on prediction accuracy but a significant impact on prediction performance.
- How are data features accessed or provided? For example, a prediction model that includes as a feature the customer's driving distance to the business location requires a service that can calculate this distance from a given address at the time of prediction (at "inference" time or at "run" time). A prediction model that includes the weather or temperature of a given day requires access to a service that can provide this information at run time. Such services need to be internally or externally provisioned, licensed and potentially paid for.

To assess model risk, relevant questions to ask by the model risk manager include the following:

- What can happen if the model acts in the worst possible way? What is the worst possible error of the model and what are the financial, business, legal or reputational impacts when the model behaves in such a way? What is the legal liability for decisions that are made based on such model behaviour?
- What if a client extracts training data or model details? For example, consider a KNN model where k is very small, e.g. 2 or 3. With just a few observations, and the prediction of the model, other observations in the training set could be identified or deduced.
- What are financial, business, legal, and reputational risks? Who assumes liability for model errors and can the liability be reduced? Are there legal implications for model errors? Are there reputational risks? The extent of such risks and the impact of model prediction error differs from use case to use case. For example, in large-scale medical diagnostics, the impacts may be much more severe than in a small pilot implementation of a dynamic pricing application.

Sources of risk that a model risk manager or model auditor should consider include the following:

- *Errors in model design or training:* Is the appropriate model type chosen? Is the model architecture appropriate to the problem and are the hyper-parameters chosen well? Did model training use the data correctly and use the correct data?
- *Errors in the runtime environment:* Are there security flaws or security "holes" in any of the software packages that are used in the runtime environment? Are

there known limitations of the packages for certain situations and do they affect the deployed model?

- *Data quality problems:* Was the training data of sufficient quality? Were data transformations and data pre-processing carried out correctly and verified?
- *Differences between training & production data ("input drift"):* Over time, the characteristics of the input data may change. For example, future customers may be different from past customers as the business's products, services, competitive position, and strategy evolve. A model that was built based on past customers' training data may no longer provide good predictive performance for current or future customers.
- *Abuse of model or misuse of outputs:* Is the model used only for the purpose it was designed for, or are its outputs also used in other ways? For example, the output of a dynamic pricing model that predicts the maximum amount a customer is willing to pay for product could also be used to offer short-term consumer credit to the customer for that product. However, the willingness to pay a particular price for an item is not the same as the ability to afford to pay a particular price or loan.
- *Adversarial attacks:* For example, consider a recommender system that predicts the probability of future movie choices on a streaming platform. Can the system be tricked into recommending age-inappropriate movies when a particular input is provided?
- *Legal risk from training data use or model results:* What are the implications of using training data for which permission to use was not obtained? Can the model be easily retrained when customers or third parties withhold or retract permission to use their data.
- *Reputational risk:* Some ML model errors are high profile and covered by national and international news outlets. This can have serious implications for the reputation of a company and its relationship with its customers.

Detailed and up-to-date documentation can help the risk manager in objectively evaluating the probability and potential impact when these risks are realized. A number of risk mitigation procedures are available to mitigate the risks, including:

- *Shadow testing* of a new model is to deploy both the old and new replacement model at the same time. Both models receive the same inputs but only the old model's predictions are used or served to the client application. This allows testing of a candidate new model on actual production data to ensure it will only go live when it is confirmed to be no worse than the existing old model. Because the new model is already deployed, switching the input and output traffic is then a simple matter.
- *Progressive rollout* (sometimes called "canary deployment") also uses both an old and a new replacement model. However, rather than shifting input and output entirely from the old to the new model, this is done progressively for increasing

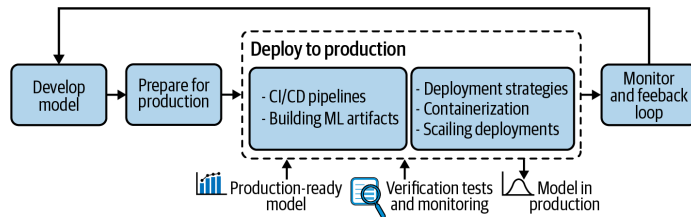
proportions of traffic. For example, initially, only 10% of customer input is routed to the new model. Then, after the model behaviour has been assessed, this is increased to 20%, etc.

- *Continuous logging and monitoring* is important to detect abnormal inputs which can signal an adversarial attack or a software malfunction. Alarms are raised and the model can be taken offline or replaced with a simpler model. Logging and monitoring is also important to detect gradual input drift, that is, changing characteristics of the input data, which indicates that the model is no longer appropriate and should be retrained.
- *Input and output checks* are quick, simple checks that ensure that input values fall within an acceptable range, are of an appropriate data type, etc. For example, a dynamic pricing model may check inputs to ensure that customer addresses are valid and in the business' geographic market. Output checks are common-sense checks that ensure the model predictions are at least sensible. For example, in a dynamic pricing application, the predicted price should be positive and within a range appropriate to the product category that is being priced.
- *Fail-over to simpler model* is the degradation of a model to a simple backup when a model failure occurs. For example, the complex neural network model in a dynamic pricing application may be unavailable because of software errors, adversarial attacks, or invalid input or invalid output. In its place, a simpler linear regression model may be used.
- *Periodic retraining* should occur whenever an input drift has been detected or a significant amount of new training data is available. This ensures that the model remains appropriate and useful for its intended purpose.

Automation tools used during the preparation for production include continuous integration and automated testing tools for quality assurance and model risk evaluation. Model registries are used during this phase to document all information about a model and document the findings of its quality assurance and risk evaluation, including input data sources and data provenance, model assumptions, required software packages, test results including explanations and bias or fairness evaluations, training and test logs, security test results, etc.

20.5.3 Deploy to Production

Deploying a model to production includes both the CI/CD (continuous integration/continuous deployment) pipelines for deploying the software that uses the model as well as building the ML model and related artifacts. The deploy to production phase is highlighted in Figure 20.11. Inputs are a production ready model and completed verification tests. The output is a model in its production environment.



Source: Treveil et al. (2020), Figure 6-1

Figure 20.11: Deploy to Production in the MLOps Lifecycle

A typical, automated CI/CD pipelines comprises at least the following steps²⁶:

1. Build model
 - (a) Build model artifacts (model code, configuration, data, trained model, environment, documentation, test code and test data)
 - (b) Archive model on model store and register model with model registry
 - (c) Basic checks for model
 - (d) Evaluate bias and interpretability
2. Deploy to test environment
 - (a) Evaluate predictive performance
 - (b) Evaluate computational performance
3. Deploy to production environment
 - (a) Limited deployment (shadow, progressive ("canary") deployment)
 - (b) Full deployment

Deployment includes considering the scalability and reliability of the model at inference time. Deployment targets (that is, server types and numbers) must be chosen that are appropriate to the expected workload. For example, an ML architect must be determine which servers should serve the model predictions, where in the world they should be located, how many there should be, and other technical considerations. Workload balancing for multiple servers must be defined and set up, for example, based on input features, request characteristics such as geographic location, or randomly. Automatic fail-over to replacement servers must be defined. This includes the ability to automatically detect server failure and to automatically re-provision replace servers. This phase must also consider how model upgrades are performed (e.g. shadow deployment, progressive deployment).

Deployment to production also includes defining and provisioning infrastructure for continuous monitoring. Three aspects should be monitored regularly. Resource mon-

²⁶ Adapted from Treveil et al. (2020) (pg. 74f)

itoring measures the infrastructure resources consumed by the model in production. This includes CPU computation time, network traffic, database and file storage space, and many related metrics. Abnormally high or low values can indicate operational problems and must be investigated. Model health checking involves assuring that the model serving software does actually accept prediction requests and provides model predictions. That is, it ensures the software works. This is different from resource monitoring which can only measure whether the server computer is busy. Finally ML metrics monitoring focuses on prediction metrics, such as input and output characteristics or their distributions, or prediction error rates or accuracy (if ground truth is available at that time or shortly after).

Automation tools required and used during this phase of the MLOps lifecycle include source code repositories such as GitHub and software integration tools such as Jenkins to build and test the model and related software applications. Also required are model registries such as MLFlow and model serving tools such as TensorFlow Serving or Flask. Finally, log data storage and log analysis software is required for monitoring.

The remainder of this section illustrates two basic deployment methods for a neural network model developed using Keras. As a first step, the model is defined and trained. The following example uses a very simple linear regression model for the Boston housing price dataset. The focus here is not on the quality of the model but on how the trained model can be deployed.

Resources

Complete implementations of all examples in this chapter are available in the following GitHub repo:

<https://github.com/jevermann/busi4720-mlops>

The project can be cloned from this URL:

<https://github.com/jevermann/busi4720-mlops.git>

As a first step, the data set is retrieved, features are created and the linear regression model is defined²⁷.

²⁷Complete implementation is available at https://github.com/jevermann/busi4720-mlops/blob/main/train_model.py

```

import keras.utils
import pandas as pd
import tensorflow as tf
import tensorflowjs as tfjs

keras.utils.set_random_seed(42)
boston_data = pd.read_csv("https://evermann.ca/busi4720/boston.csv")

boston_features = boston_data[['rm', 'tax', 'age']]
boston_labels = boston_data['medv']

# Linear regression model
norm_boston_model=keras.models.Sequential([
    keras.layers.Input(shape=(3,), dtype=tf.float32),
    keras.layers.Dense(1, activation=None) ])

```

Next, the model is trained and saved in three different ways, for use in Keras, for use in TensorFlow serving, and for use in TensorFlowJS. Each of these require a different model package format.

```

stop_callback = keras.callbacks.EarlyStopping()
norm_boston_model.compile(
    loss = tf.keras.losses.MeanSquaredError())
norm_boston_model.fit(
    boston_features, boston_labels,
    epochs=100, validation_split=0.33,
    callbacks=[stop_callback])

# Save model for use in Keras
norm_boston_model.save('norm.boston.model.trained.save')
# Export model for use in TF Serving
norm_boston_model.export('norm.boston.model.trained.export')
# Convert model for use in TFJS
tfjs.converters.save_keras_model(norm_boston_model,
    'norm.boston.model.trained.tjfs')

```

Deployment as a Flask Microservice

The following code blocks illustrate how a trained model may be deployed as a micro-service using the Flask WSGI server. Flask is a Python package that functions as a web server, accepting requests from client applications over the network and sending an appropriate response. In this example, client applications send the input data for the prediction model as a JSON object in the web request, and the Flask service responds with a JSON object that contains the prediction and other information²⁸.

First, the saved model is loaded. The `predict()` function accepts the inputs and

²⁸Complete file is available on https://github.com/jevermann/busi4720-mlops/blob/main/flask_deploy.py

provides suitable output (assuming that inputs are single observations, it returns the first and only target of the first and only element of a batch).

```
import keras
import flask
from flask import request
import pandas as pd

# Load the trained model
norm_boston_model = keras.saving. \
    load_model('norm.boston.model.trained.save')

# A predict function for the model
def predict(inputs):
    return norm_boston_model.predict_on_batch(inputs)[0][0]
```

Setting up the Flask web server is simple. The app defines the Flask application and the decoration `@app.route(...)` indicates that the `predict_json()` function should be called when a client requests the `/predict_json` URL using the POST method on the server. The `predict_json()` function accepts the input data from the request, and creates a suitable Pandas dataframe. It calls the `predict()` function and packages the resulting prediction as a JSON object to be returned to the client.

```
app = flask.Flask(__name__)

# Define the URL handler:
@app.route("/predict_json", methods=["POST"])
def predict_json():
    reply = {}
    # TODO: Input checking goes here
    # TODO: Input logging goes here
    inputs = pd.DataFrame.from_dict(request.json).transpose()
    prediction = predict(inputs)
    # TODO: Output checking goes here
    # TODO: Output logging goes here
    reply["prediction"] = str(prediction)
    reply["success"] = True
    return flask.jsonify(reply)

# Run the server app
app.run()
```

To access this microservice application when it is running, a simple web request can be used from the Bash command line²⁹. The curl program sends web requests and receives and prints the responses, the `-X` option specifies the request method, the `-H` option specifies request headers, `-d` specifies the data to be sent and the final argument is the web server address and URL.

²⁹Complete file is available on https://github.com/jevermann/busi4720-mlops/blob/main/json_demo.sh.

```
curl -X POST \
-H "Content-Type: application/json" \
--data '[6, 250, 66.5]' \
http://localhost:5000/predict_json
```

To show how such a service may be used from a web-based application, the following code blocks define a simple HTML form³⁰. The form provides input fields for the user to enter the input feature values, and in the document header defines a short JavaScript script that calls the Flask microservice to provide the predicted house price.

```
<!DOCTYPE html>
<html lang="en">
<head>
<meta charset="UTF-8">
<title>Boston Housing Data Prediction Service</title>
<script>
async function predict() {
    // Get the values from the text inputs
    const rooms = parseFloat(document.getElementById('rooms').value);
    const tax = parseFloat(document.getElementById('tax').value);
    const age = parseFloat(document.getElementById('age').value);
    // Make a POST request to the server
    const response = await fetch('/predict_json', {
        method: 'POST',
        headers: { 'Content-Type': 'application/json' },
        body: JSON.stringify([rooms, tax, age]);
    });
    // Parse the JSON response
    const result = await response.json();
    // Display the result
    document.getElementById('output-div').textContent
        = result.prediction;
}
</script>
</head>
```

The body of the HTML document is the HTML form with its input elements. When the "submit" button is pressed, the `predict()` function defined in the document header in the previous HTML code block is executed.

³⁰Complete file is available on https://github.com/jevermann/busi4720-mlops/blob/main/predict_form_async.html.

```

<body>
  <h1>Boston Housing Data Inputs</h1>
  <form onsubmit="event.preventDefault(); predict();">
    <p>
      <label for="rooms">Number of Rooms</label>
      <input name="rooms" id="rooms" required>
    </p>
    <p>
      <label for="tax">Tax Rate per 10,000</label>
      <input name="tax" id="tax" required>
    </p>
    <p>
      <label for="age">Prop bldg older than 1940</label>
      <input name="age" id="age" required>
    </p>
    <input type="submit" value="Submit">
  </form>
  <p>Prediction is: <div id="output-div">...</div></p>
</body>
</html>

```

The Flask application defined here is typically packaged in container (using Docker or other container technology). One or more instances of such a container are then executed on container servers, depending on the required capacity. Container deployments to servers are typically automatically performed using software such Kubernetes³¹, which is an open-source container orchestration system for automating software deployment, scaling, and management.

Deployment as a JavaScript Model

Another way to deploy an ML model is to embed it in a browser-based web application. This deployment has the advantage that the organization does not need to provide computational resources to run the model at inference time, and the prediction request and response need not be sent across a network connection. On the other hand, this deployment is suitable only for small models and only for models that can be made available to application users as organization loses control over the model and must treat its results as insecure. Examples where such models may be useful could be object recognition tasks in a web browser, speech translation or transcription in a web video conferencing application, etc.

The following HTML code block³² is a slightly altered version of the web form in the Flask example above. Instead of requesting a prediction from the Flask service, the JavaScript script loads the model directly using the `tf.LoadLayersModel()` function provided by the TensorFlowJS framework and calls its `predict()` function. The document body that defines the form and its input fields is identical to the one in the code block above.

³¹<https://kubernetes.io/>

³²Complete file is available on https://github.com/jevermann/busi4720-mlops/blob/main/tjfs_demo.html.

```

<!DOCTYPE html>
<html>
<head>
<script src="https://cdn.jsdelivr.net/npm/\
@tensorflow/tfjs@latest/dist/tf.min.js"></script>
<script>
async function predict() {
  // Load the model
  const model = await \
    tf.loadLayersModel('https://raw.githubusercontent.com/jevermann/busi4720-mlops/main/model.json');
  // Get the values from the text inputs
  const rooms = parseFloat(document.getElementById('rooms').value);
  const tax = parseFloat(document.getElementById('tax').value);
  const age = parseFloat(document.getElementById('age').value);
  // Package the values into a Tensor
  const inputs = tf.tensor2d([rooms, tax, age], [1, 3]);
  // Get the prediction from the model
  document.getElementById('output-div').innerText =
    model.predict(inputs).dataSync();
}
</script>
</head>

```

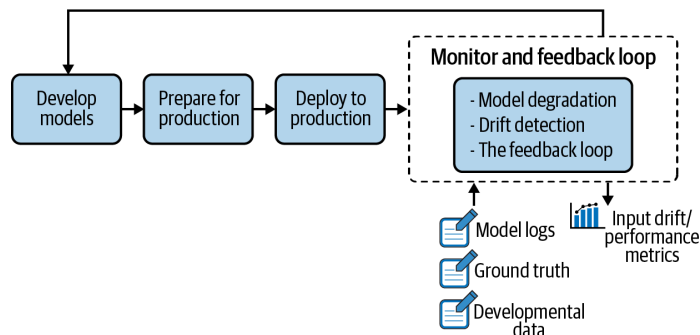
20.5.4 Monitoring and Feedback

The monitoring and feedback phase of the MLOps lifecycle, highlighted in Figure 20.12, is important to ensure that the model performance does not diminish or degrade over time. It also detects any input drift, that is, systematic changes in the characteristics or frequency distributions of the input values, and provides a feedback loop for subject matter experts.

Inputs to this phase are the model logs, the ground truth, and developmental data for the deployed model. Ground truth denotes the actual, true value for the predicted target. Output of this phase are metrics that quantify input drift and model performance.

The feedback loop is important in that it triggers model retraining when required. The input drift and performance metrics are evaluated and assessed by the subject matter expert with reference to the relevant business key performance indicators. However, model retraining is not automatically triggered but depends on a number of considerations, for example:

- *Domain changes*: This indicates a shift or change in business requirements. For example, rather than extracting the maximum price for each item from a customer in a dynamic pricing model, the business is now also interested in maximizing the purchase of related products. Therefore, the existing prediction model may no longer be suitable or useful.
- *Training cost*: Model training, especially for complex models or large training data sets, is time consuming and computationally costly. Organizations must



Source: Treveil et al. (2020), Figure 7-1

Figure 20.12: Monitoring and Feedback in the MLOps Lifecycle

consider these costs and the potential benefits or a more accurate prediction.

- *Model performance*: Not every change in prediction accuracy or prediction error translates into a significant negative business impact. Some changes may be tolerable, in particular in light of the potential training time and costs for a new model version.
- *Ground truth availability*: While continuous monitoring and logging can provide additional input data for training, it does not also provide the true target values. For example, while a dynamic pricing prediction service captures the input feature values, and whether a customer purchases a product offered at the predicted price, the true maximum price that a customer is willing to pay is not known.

Ground truth availability is a significant problem in many ML applications for at least the following three reasons:

- *Ground truth is not immediately or imminently available*. Consider the example of the dynamic pricing service that predicts the maximum amount a customer is willing to pay. The true maximum will never be known. Proxy variables can be used, for example whether a customer purchases a product at a given price, but these remain approximations. An example where ground truth is available late is loan application prediction, where the model predicts loan repayment or default of a customer. Loan repayment can only be fully captured at the conclusion of the loan duration, which may be months or even years after the prediction was made.
- *Ground truth and prediction are decoupled*. Difficulties in capturing the ground truth can arise when different software applications are involved. For example, the dynamic pricing prediction model may not include a customer's identification as an input so that this information is not logged. This makes it difficult to later identify whether a purchase occurred for a particular price prediction.
- *Ground truth not available for all classes*. Consider a fraud detection or predic-

tion service. Predicted instances of fraud ("positive predictions") may be investigated and be shown to be either true or false. that is, true and false positives will be known. However, not all negative predictions can or will be investigated because of their large number. Therefore, true negatives and false negatives will not be identified.

Input drift, that is, systematic changes to the frequency distributions of input values, may be caused by a non-stationary environment, for example, customers or markets changing over time, or by selection bias induced by the model itself. For example, a dynamic pricing application may drive away price conscious customers from the web store, changing the characteristics of the input values.

There are two main methods for input drift detection. The first method uses univariate statistical tests, for example, the χ^2 or Kolmogorov-Smirnov tests^{33 34} to test whether two samples of input values are drawn from the same probability distribution. While these tests are simple and quickly carried out, they neglect the multi-variate distribution characteristics of the input data.

The second approach is called the "*domain classifier approach*". Conceptually, the old and new input values are considered as two classes ("domains"), and a classifier is trained on the input values themselves to predict whether an input observation belongs to the old class or the new class. If such a classifier can be trained to yield a classification accuracy better than chance, the old and new input data sets can be distinguished and must be considered different.

The monitoring and feedback phase of the MLOps lifecycle requires software tools for logging of inputs, predictions, model explanations, and actions taken by the consumer of the prediction (e.g. the customer). This phase also requires access to model stores and online evaluation support. The latter tools allow for A/B testing or shadow testing, that is, to have two or more models in production and learn which one performs better based on live data.

To illustrate basic logging in Python, the following code blocks extend the Flask microservice application introduced above³⁵. This example can illustrate only the most basic notions of logging events to a log file for later analysis. It uses the built-in, default logging package for Python.

The first code fragment sets up a logger to log the web requests to a file. A logger is characterized by its name and log level. Typical log levels, across many logging frameworks in most programming languages, are DEBUG, INFO, WARN, ERROR and CRITICAL, which are ordered in increasing order of severity. For example, DEBUG logging can be used to log a multitude of events that are useful to know about when creating a software application. However, DEBUG logging is generally not needed for applications in production. INFO logging captures events that are useful to determine

³³https://en.wikipedia.org/wiki/Chi-squared_test

³⁴https://en.wikipedia.org/wiki/Kolmogorov-Smirnov_test

³⁵Complete file is available on https://github.com/jevermann/busi4720-mlops/blob/main/flask_deploy_logging.py.

the normal functioning of an application whereas `WARN` events in a log indicate conditions that are abnormal, but have not led to errors. Finally, `ERROR` events indicate errors in a software application but the application continues to work, while `CRITICAL` log events indicate errors that have led to the halting of a software application. The code below sets the logging level to `INFO` as routine events are to be captured.

A file handler is added to the logger for writing the events to a file, rather than printing them on the console. As an alternative, a `RotatingFileHandler` may be used. This handler automatically limits the log file size and rotates logs when a file size exceeds the limit. In the example below, the 5 most recent logs are kept, while earlier log information is discarded.

```
import logging.handlers

# Set up the logger:
req_logger=logging.getLogger(model_name+'.requests')
req_logger.setLevel(logging.INFO)
req_logger.addHandler(
    logging.FileHandler(
        model_name+'.requests.log'))
# req_logger.addHandler(
#     logging.handlers.RotatingFileHandler(
#         model_name+'.requests.log',
#         maxBytes=1000000,
#         backupCount=5))
```

With the request logger defined, the following code fragment shows how it can be used. The code block modified the `predict_json()` and the `predict_form()` functions to log the web request, together with the model name, the current time, and the remote internet address (the address of the web client or browser). Logging is done simply by calling the `info()` method of the request logger and providing the message to be logged as well as the arguments to be inserted into that message.

```
# Use the logger:
@app.route("/predict_json", methods=["POST"])
def predict_json():
    req_logger.info('%s TIME %s IP %s JSON %s',
                    model_name,
                    time.ctime(),
                    request.remote_addr,
                    request.json)
    ...
def predict_form():
    req_logger.info('%s TIME %s IP %s FORM %s',
                    model_name,
                    time.ctime(),
                    request.remote_addr,
                    request.form)
    ...
```

Hands-On Exercise

1. Download the complete file from [GitHub](#).
2. Define a second logger that writes to a different log file
 - You do not need to rotate this log file
 - The definition of the second logger is analogous to that of the request logger
3. Add logging to the `predict_json()` and the `predict_form()` functions to capture the time, the three input values, and the prediction outcome in the log.
 - Replace the `# TODO: Output logging goes here` comments with your code
 - To make the log easy to analyze, write the information in CSV format. Make sure you quote the fields that need quoting.

20.6 ML Governance

In general, governance provides control, oversight, guidance, and makes strategically important decisions. ML governance in particular provides oversight, control, and directions to ML operations, and makes important strategic decisions around ML use in an organization. Governance often asks critical questions of management and operations to ensure that risks are appropriately managed and business value is realized or organizational goals are achieved. Figure 20.13 shows an 8-step process model for ML governance³⁶.

1. *Understand ML Use Cases.* This step prompts the organization to understand the extent to which it uses ML models and for what purposes. Typical questions to ask (and to answer!) are the following:
 - Who are the consumers of the model outputs? Some models may be developed for internal consumption and decision making, while others may be customer facing.
 - What regulations and legal constraints apply to each use case or model? Different industries and geographic or jurisdictional areas have different constraints. Organizations must identify the users of their models and the laws and regulations that apply.
 - What are the legal, financial, reputational risks of prediction errors? Answering this question ensures that effective model risk management is in place, as described above.
 - What is need for explainability or interpretability? Not all models need to be interpretable, and different models or model use cases may require different types of interpretability.

³⁶The figure and the material in this section are adapted from Treveil et al. (2020).

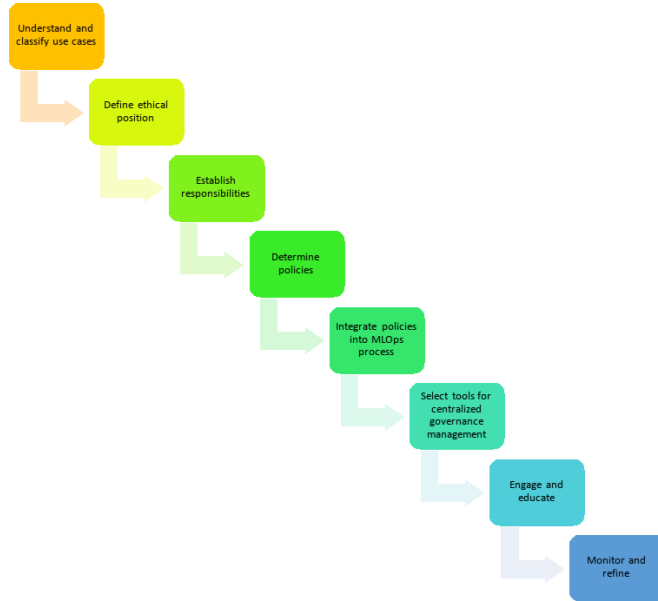


Figure 20.13: ML Governance Phases (after Treveil et al. (2020), Chapter 8)

- What are the availability requirements for the various models? Not every model must provide 5-nines uptime, that is, be available 99.999% of the time. Ensuring availability is generally costly.
- What are the model lifetimes and likely rates of model deterioration? Knowing even approximate answers to this question can inform how fast or responsive the MLOps processes need to be.

2. *Define the Ethical Position.* This step guides how ML models will be used (or not used) by an organization. It examines the following questions:

- How important are aspects like equality, privacy, human rights, democracy, bias? Answers to this question can, for example, inform the features that are used or excluded from prediction models, or what data is stored and logged from models in production.
- How transparent should decision making be to the customers of the prediction models? Answers can inform the use of interpretable ML methods or the type of models deployed by the organization.
- What level of responsibility for errors will or should the business assume? For some errors, the business or organization may accept financial responsibilities, for others it may try to deflect this responsibility by appropriate legal instruments (e.g. terms-of-service or licenses to use).

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Tasks	Business stakeholders	Business analysis/citizen DS	Data scientists	Risk/audit	Data ops	Production/exploitation	Resources admin/architect
Identification	A/R	C		I			
Data preparation	C	A/R	C				
Data modeling	C	A	R				
Model acceptance	I	C	C	A/R			
Productionalization		C	A/R	I	C		
Capitalization			R		R		A
Integration to external systems					A/R		
Global orchestration		C			R	A	
User acceptance tests	A/R	R	C		I		
Deployments					R	A	I
Monitoring	I	C				A/R	I

A: accountable R: responsible C: consulted I: informed

Source: Treveil et al. (2020), Figure 8-4

Figure 20.14: RACI Matrix for ML Governance

- What is the potential for deception, manipulation, exploitation of model users? For example, a recommendation system that is used to increase sales may lead some users to purchase products or services they cannot afford, leading to financial hardship. As another example, consider recommendation systems for news articles that may lead users to biased or limited coverage.
3. *Establish Responsibilities.* This step essentially defines "who will do what?". A frequently used way to define responsibilities is through the use of a "RACI" matrix as in Figure 20.14. RACI stands for "responsible", "accountable", "consulted", and "informed". This step of ML governance covers the following points:

- Responsibilities must be defined at the strategic, tactical, and operational level. For example, who must be involved in model training (operational), who is allowed to sign-off on model risk (tactical) and who determines business goals or KPIs for a model (strategic)?
- Senior management sponsorship is important for the MLOps lifecycle. Without senior management recognizing the importance of ML models to the organization, and providing adequate resources and guidance for their development and use, ML and MLOps cannot function.
- ML governance should be integrated into existing governance mechanisms. For example, organizations typically have IT governance mechanisms and a set of IT related policies and procedures. ML governance can extend these and define whether and how they apply to the MLOps lifecycle.

4. *Define Policies.* This step answers the question "how will we do that?". Policies are formal and explicit rules that govern the details of what acceptable performance of the following issues means and how acceptable performance levels are achieved and documented. The following are the major MLOps aspects that require policies and some example questions to be answered by such policies:

- Reproducibility and traceability: How is this achieved, measured, and verified? What are acceptable levels?
- Auditability and documentation: How much documentation and in what format is required? Where is the documentation maintained and who can access it?
- Sign-off between stages: Who has authority to, for example, select a model and move it to testing or deployment?
- Model verification: What verification tests must be performed? What are acceptable results? Who determines acceptable results? How are verification results documented?
- Model explainability: What types of explainability are required for each model? Who is allowed to define such requirements? How can they be tested and documented?
- Model bias and bias testing: When and how must model bias be tested? What are acceptable levels of bias or fairness? Who determines them? How is bias testing documented?
- Model deployment mechanisms: Who has the authority to decide the choice of deployment mechanism and how are the resulting infrastructure requirements and model risk determined? How are decisions and their rationale to be documented?
- Model monitoring: At what intervals is model performance monitored? Which metrics are recorded? How long are such records kept and maintained? Who has access to such records for analysis? What alarms are defined to identify model problems? Who must react how quickly to such alarms?
- Data quality and data compliance: What are acceptable levels of data quality? How is it ensured, tested, and documented? Who is responsible for data compliance with legal and regulatory constraints? How is compliance demonstrated?

As noted, existing governance mechanisms, such as an IT use policy or an IT development or DevOps policy, should be applied when relevant.

5. *Integrate Policies into MLOps Process.* This step ensures that the relevant policies are actually applied and followed throughout the phases of the MLOps life-cycle. In particular, integration involves the following steps:

- Formalize and automate MLOps processes: Automation of processes can ensure that the actions and decisions specified in policies are applied reliably and consistently. As far as possible, the activities required by a policy should be tool supported or automated.
 - Define controls: This step translates policy requirements into actionable or specific constraints and determines how control effectiveness can be measured. For example, if a policy specifies that a model must be globally explainable in terms of feature importance, an automated tool should prevent a model being deployed to production that does not have associated explanation test results in the model repository. A control for this may specify exactly what the model repository must contain. This also indicates how control effectiveness can be tested: All models in production should have the required explanation test results.
 - Define monitoring of controls: When and how frequently is control effectiveness monitored? Who performs such monitoring and who is responsible for actions in case of violations?
6. *Implement Governance Tools.* As one of the core principles of MLOps is automation, it is unsurprising that ML governance mechanisms should also be automated and supported by appropriate software tools. This involves the following aspects:
- Automate controls: As indicated above, controls should be automated in the software tools that form part of the MLOps tooling, such as model repositories, ML pipelines and ML workflow orchestration tools.
 - Logging of control violations: Once controls are defined, violations should be monitored and logged. This ensures accountability and auditability but is also useful for determining how MLOps processes or policies can be improved in the future.
 - Auditing of control effectiveness: Policy compliance and control effectiveness should be periodically audited by an independent auditor, that is, staff outside the MLOps process. Examples might be staff of the chief risk officer (CRO) or of the chief information officer (CIO) of the organization.
 - Policy and procedure maintenance: Policy repositories can support periodic policy and procedure evaluation, assessment and updates when required. They support policy dissemination and version management.
7. *Engage and Educate.* Policies and procedures are only effective if they are known and followed by those who they apply to. This step in the governance process includes the following aspects:
- Communicate: Organizations have many different formal, semi-formal, and informal ways to communicate policies and procedures.
 - Awareness: While communication makes policies and procedures available, awareness requires that MLOps participants consider the applicable

policies throughout their daily work.

- Training: Different training methods, online or in-person, individual or in groups, must be made available that cover both technical MLOps aspects as well as applicable policies, and how to best implement or comply with policies.
- Buy-in & commitment: Successful MLOps and ML governance requires the buy-in and commitment to ML governance and efficient MLOps of all stakeholders and participants, at all levels of the organizations.
- Culture: Successful MLOps is a cultural issue as much or even more so than a technical one.

8. *Monitor and Refine*. Just like deployed models must be continually monitored in the MLOps lifecycle, so must deployed policies and controls be continually monitored in ML governance. This includes:

- Evaluating risk exposure: Governance, through the deployed policies, is designed to manage risk. An organization must periodically assess the actual risk exposure versus acceptable levels of risk and must, if necessary, adapt its governance mechanisms such as responsibilities, policies, or specific controls.
- Evaluating policy adequacy: This examines whether policies are appropriate to control risk and ensure efficiency of the MLOps process.
- Evaluating control effectiveness: As noted above, control effectiveness must be continually monitored and evaluated. If necessary, controls must be strengthened.
- Evaluating MLOps process performance: The overall efficiency and effectiveness of the MLOps process must be evaluated. This asks questions such as: "how long does it take to move a new model to production?", "how long does it take to recover from a production model problem?", "how often are there significant problems when moving a candidate model through testing?". Recall that governance mechanisms are not only intended to manage risk but also to ensure efficiency. When process performance does not meet expectations, processes and policies may need to be adapted.

20.7 Review Questions

Introduction

1. Why is the integration of a trained model into a production environment important, and what challenges can arise during this process?
2. Provide an example of how a predictive model might be used in a business context and the potential risks involved.
3. What are the three main purposes of MLOps?

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4. Why is the constant change in data a significant challenge for organizations using predictive models?
5. Describe the composition of business analytics teams and the issues that may arise within such teams.
6. What are the main principles around moving predictive analytics models into production according to MLOps?
7. Define "infrastructure as code" and its importance in MLOps.
8. Describe how MLOps intersects with machine learning, software engineering, and data engineering.

MLOps Lifecycle Overview

9. Describe the two lifecycles that are combined in the MLOps lifecycle.
10. How does the DevOps approach address the issues found in early software development practices?
11. Explain the rationale behind combining the model development lifecycle and the DevOps lifecycle.
12. How does the combined MLOps lifecycle address the integration of machine learning models into complex software applications?
13. Using the example of a dynamic pricing web-store application, illustrate the steps involved in the MLOps lifecycle.
14. Discuss the importance of formalized and automated processes in managing the MLOps lifecycle efficiently.

MLOps Roles and Requirements

15. Describe the role of subject matter experts in the MLOps lifecycle. What are the requirements for subject matter experts to perform their role efficiently?
16. Explain the responsibilities of data scientists in the MLOps process.
17. How do data engineers support data scientists, and what are their main responsibilities?
18. What automated tools do data scientists and data engineers require to perform their tasks efficiently?
19. Discuss the role of data engineers in the MLOps lifecycle.
20. Define the role of software engineers in integrating machine learning models. Describe the requirements of software engineers for effective collaboration and code management.
21. What are the responsibilities of DevOps engineers in the MLOps lifecycle?
22. Who are model risk managers and model auditors, and what are their primary responsibilities? What tools do model risk managers require?
23. Describe the roles of ML engineers and ML architects in the MLOps lifecycle. Why is it important for ML engineers to quickly assess and adjust infrastructure capacities?

MLOps Tooling

24. What are CI/CD tools, and how do they support the MLOps lifecycle?

25. Define feature stores and their importance in managing data for ML models.
26. Discuss the role of model registries in the MLOps lifecycle.
27. What is the purpose of ML metadata stores, and how do they support MLOps?
28. Explain the importance of model monitoring tools for deployed models.

MLOps Lifecycle Phases

29. Describe the main phases of the simplified MLOps lifecycle.
30. What are the main activities in and the outputs of the model development phase of the MLOps lifecycle?
31. List some important questions related to data management that should be addressed during the model development phase.
32. What are some key considerations for ensuring model fairness and mitigating bias during model development?
33. What are the main tasks in and the inputs and outputs of the phase of preparing a model for production?
34. What are the key considerations when selecting a runtime environment for a model?
35. What are some key technical questions to consider when preparing a model for production?
36. Describe the potential risks associated with deploying a machine learning model and how these risks can be assessed.
37. List and explain some risk mitigation procedures that can be employed during the preparation for production phase.
38. What are the main steps in and the inputs to deploying a model to production?
39. Describe the components of a typical automated CI/CD pipeline for deploying machine learning models.
40. What are the key considerations for ensuring scalability and reliability of model inference during deployment?
41. Explain the importance of continuous monitoring in the deployment phase and what aspects should be monitored.
42. Explain how a Flask microservice can be used to deploy a trained model.
43. How can a neural network model be deployed in a browser-based web application using TensorFlowJS?
44. Explain the concept of input drift and its potential impact on model performance.
45. What are some aspects to consider when triggering model retraining?
46. Discuss the challenges associated with ground truth availability and how they impact the monitoring and feedback phase.
47. Describe two main methods for detecting input drift.

ML Governance

48. What is the role of ML governance in the context of ML operations?
49. Why is it important for an organization to understand its ML use cases?
50. What are some critical questions to ask when understanding ML use cases?

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51. What are the potential consequences of not identifying the legal and regulatory constraints for ML models?
52. Discuss some ethical considerations that organizations must address when using ML models.
53. How can transparency in decision-making affect the deployment of ML models?
54. Why is it important for an organization to consider the potential for deception or manipulation by ML models?
55. Explain the concept of a RACI matrix and its role in ML governance.
56. Why is senior management sponsorship important for the MLOps lifecycle?
57. What are the key aspects of MLOps that require defined policies?
58. Give examples of questions that need to be answered by policies related to model verification and monitoring.
59. Provide examples of questions that policies should answer for reproducibility and traceability.
60. Why is it important to formalize and automate MLOps processes?
61. How can governance tools automate controls in the MLOps process?
62. Explain the importance of logging control violations and auditing control effectiveness.
63. List the aspects involved in engaging and educating MLOps participants.
64. What is the importance of monitoring and refining policies and controls in ML governance?
65. How can policy repositories support policy and procedure maintenance?

Appendix A

Installing and Using a Virtual Machine

A.1 Introduction

A virtual machine provides a "computer within the computer". That is, it is a complete "*guest*" computer system with its own hard drive space that can run as an application on a "*host*" computer system. This makes it ideal to deliver a fixed environment with software packages and data sets fully installed and configured.

A virtual machine is specific to a particular type of computer hardware (processor chip/CPU) and the guest system hardware should be the same as the host system hardware. Hence, when choosing how to proceed, it is important to determine the real computer's hardware. As of this writing, there are two major processor chip/CPU types in use.

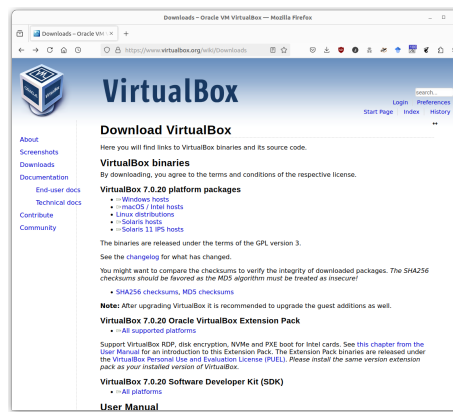
- *Intel and the fully compatible AMD chips* are used primarily with the Windows operating system. However, they were also used by Apple Mac computers until approximately 2021.
- *ARM chips* are used in current Apple Mac computers (since approximately 2021, using the Apple M1, M2, or M3 processors/CPU), but are also used in a number of recent Windows laptops (since approximately 2023, using the Qualcomm processors/CPU).

The virtual machine for this course is available for Intel/AMD processors running Windows or MacOS, as well as for ARM processors running MacOS. ARM processors running Windows have not been tested.

A.2 VirtualBox on Windows

The virtual machine for use on Intel and AMD processors (CPUs) uses the Oracle VirtualBox virtual machine software application. Oracle VirtualBox is free and open-source software that is available at no cost. It comes packaged as a virtual box appliance for import into the VirtualBox software. To install the virtual machine, follow these steps:

1. Ensure you have approximately 80GB of hard drive space available (30GB to download the virtual box appliance file, and 50GB to install it).
2. Download Oracle VirtualBox from its [website](http://www.virtualbox.org/wiki/Downloads). Choose the version for your operating system (Windows or MacOS). The virtual machine was developed with VirtualBox version 6.1 but should work with newer versions of VirtualBox.

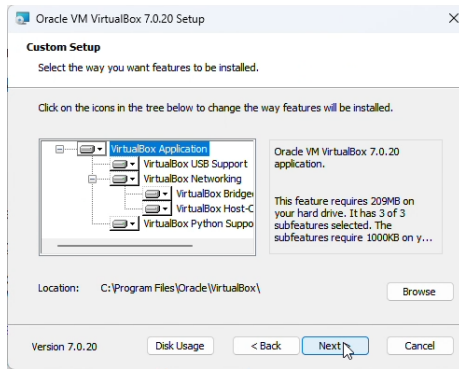


3. Follow the [general installation instructions](#) or the [detailed instructions for Windows](#) or the [detailed instructions for MacOS](#). The following screen shots guide you through the installation of VirtualBox on Windows:

- The setup wizard guides you through the setup process. You should confirm default choices:



- Confirm the default features to install:



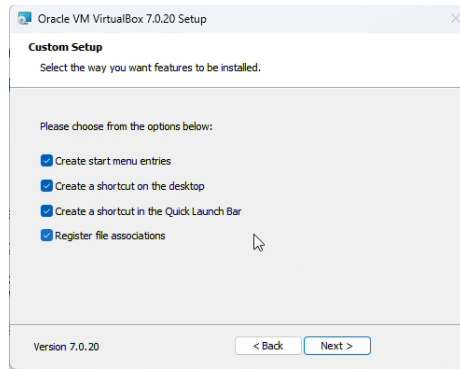
- Confirm the warning about temporary network disconnect:



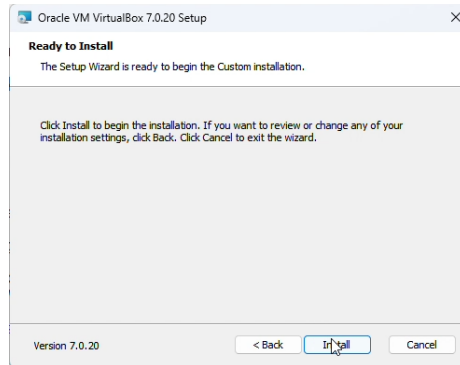
- Ignore the warning about missing Python dependencies:



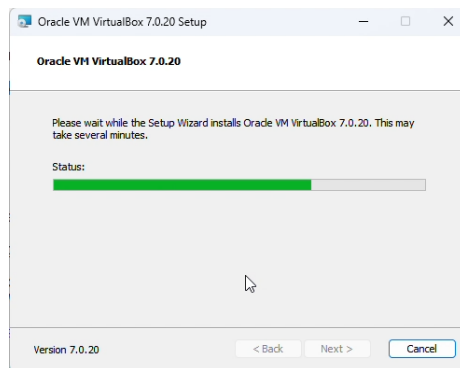
- Select which menu entries and shortcuts to create according to your preferences:



- Confirm the installation:



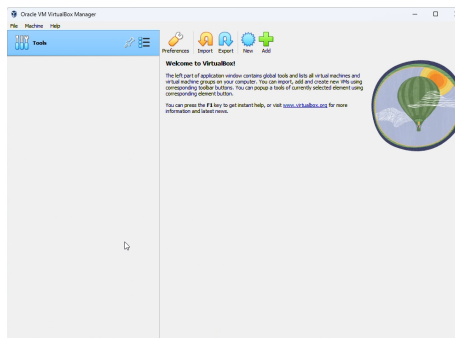
- Wait for the installation to finish:



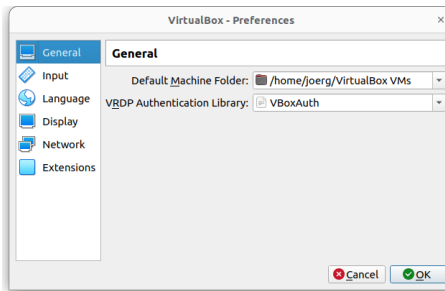
- Finish the installation, start VirtualBox, and exit the setup wizard:



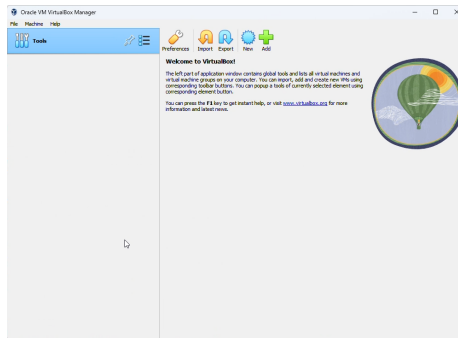
- You will see the VirtualBox Manager screen:



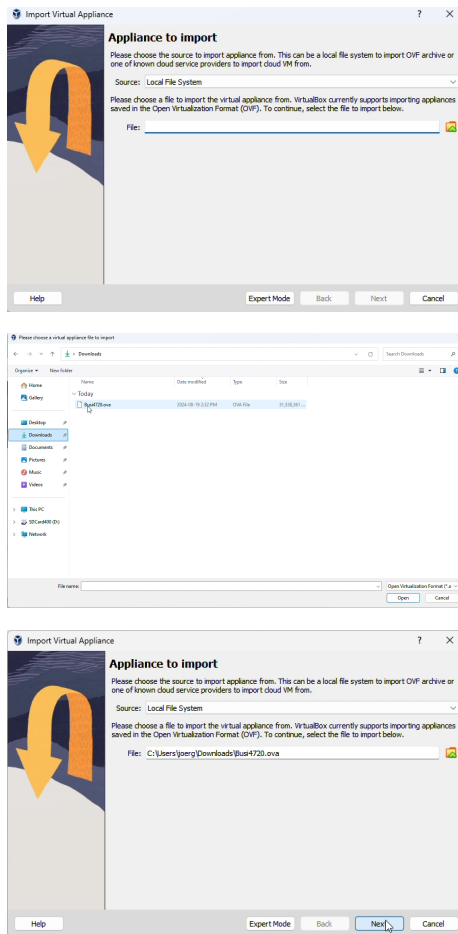
- Press the "Preferences" button. In the general preferences, specify the location for VirtualBox to store the virtual machines. Ensure at least 50GB of hard drive space is available in that location:



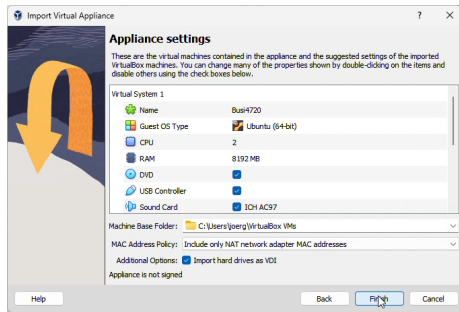
- An overview of the VirtualBox user interface is found in [chapter 1 of the user manual](#).
4. Download the [virtual appliance file](#). *Warning:* This is a 30GB file and will take some time to download.
 5. Import the virtual appliance into the VirtualBox software.
 - On the VirtualBox Manager main screen, select "Import"



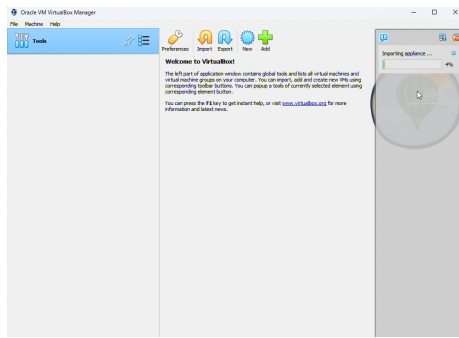
- Select the virtual appliance file you have downloaded. This is most likely in your "Downloads" folder



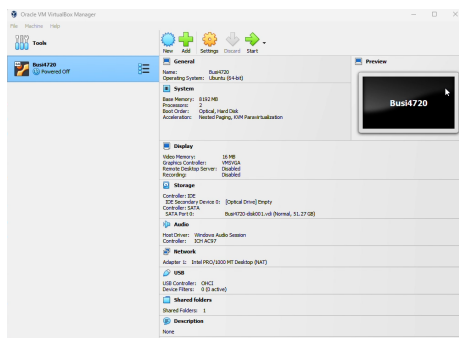
- Confirm the appliance settings. At this point, you may also choose a different folder than the default one for storing the virtual machine:



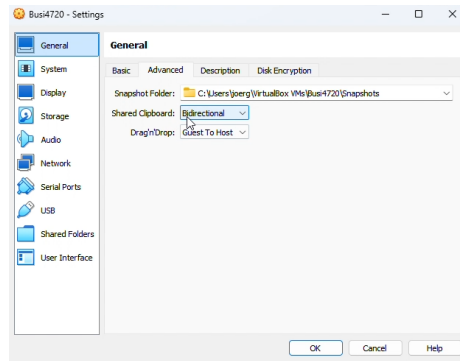
- Wait for the import to complete. This may take a few minutes.



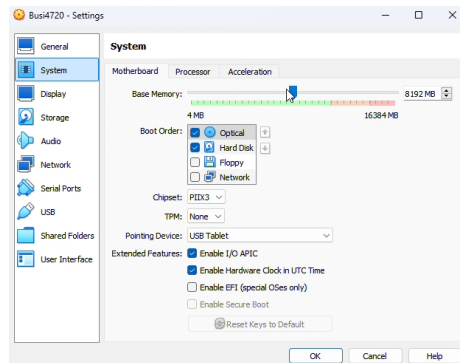
- When the import is complete, the virtual machine settings will be displayed. Press the "Settings" button to change settings for the virtual machine.



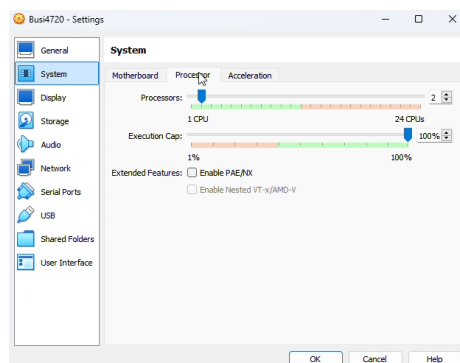
- In the General settings, you may choose whether to share the clipboard for copy/paste between guest and host, and whether to enable drag-and-drop operations between guest and host.



- In the Systems settings, you may choose how much memory to allocate to the virtual machine. You should allocate around 8000 MB, but ensure that enough memory is left over for the host computer to work smoothly.

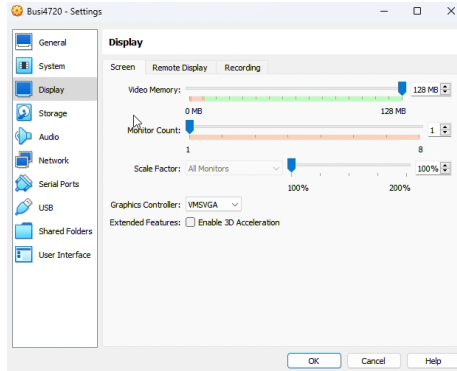


- In the Processor tab of the Systems settings, you may choose how many processor chips/CPUs to allocate to the virtual machine. Ensure that enough are left over for the host computer to work smoothly.

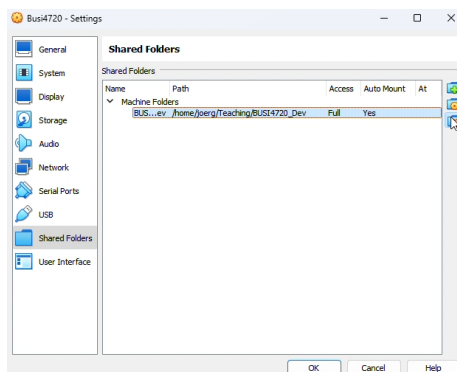


- In the Display settings, you should allocate all the maximum video memory to the virtual machine. You may also choose a scaling factor. This may be useful if you are working on a very high resolution monitor like an Apple

Macbook Retina display.

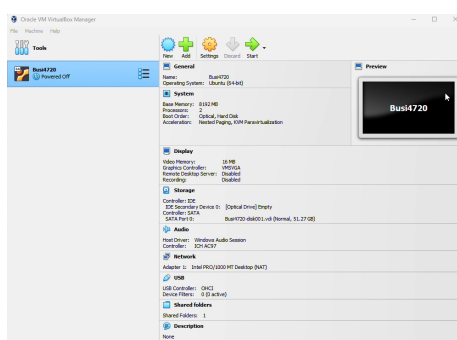


- In the Shared Folders settings, you should remove any existing shared folders.



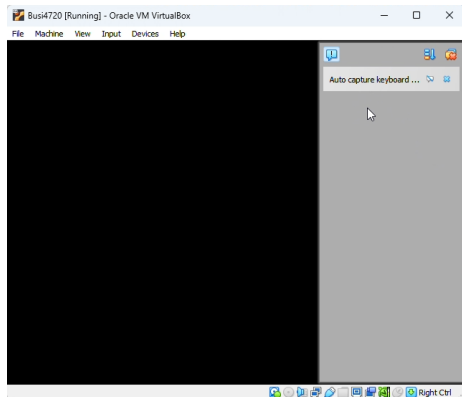
6. Start the virtual machine.

- Press the "Start" button on the virtual machine settings overview.

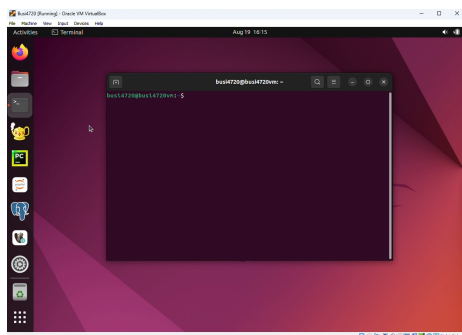


- The virtual machine will start and notify you about keyboard and mouse integration. This means your keyboard and mouse inputs will automatically be forwarded from the host to guest system. You can dismiss these

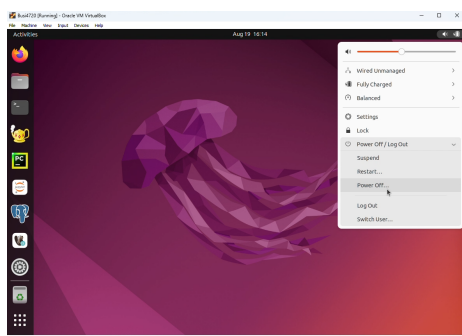
messages



- Once the virtual machine has started, you will see the desktop user interface. Popular software applications are accessible via the dock on the left side of the screen. The most important application for working with this book is the Terminal window application.



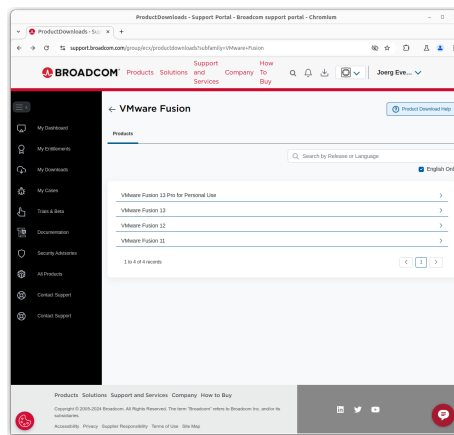
- To shut down the virtual machine, select "Power Off..." from the system menu in the top right and shut down the machine.



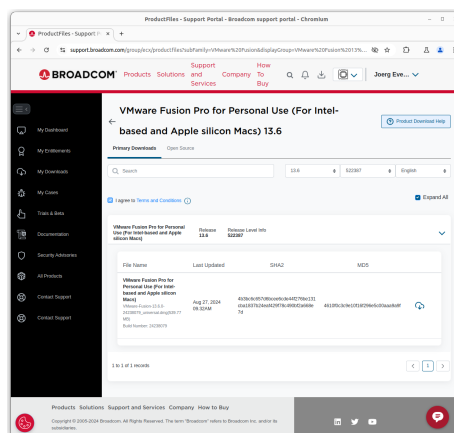
A.3 VMWare Fusion on MacOS

The virtual machine for use on ARM processors (CPUs) uses the VMWare Fusion Pro virtual machine software. VMWare was acquired by Broadcom in 2023. While VMWare Fusion is proprietary software, Broadcom makes it available at no cost for personal use. To install the virtual machine, follow these steps:

1. Ensure you have approximately 70GB of hard drive space available (20GB to download the virtual machine file, and 50GB to install it).
2. Download VMWare Fusion Pro from its [website](#). You may need to register for a free Broadcom account and login.
 - Choose the latest version of VMWare Fusion Pro for Personal Use. The virtual machine was created with version 13.5.2.



- Tick the box to agree to terms and conditions and press the download button next to the VMWare Fusion product.

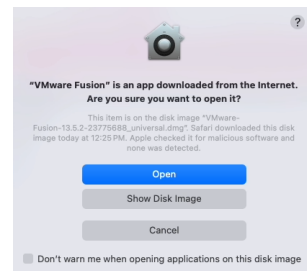


3. Install VMWare Fusion Pro

- Once the download is complete, double-click the downloaded file. You will see the VMWare Fusion installer. Double-click it to begin the installation.



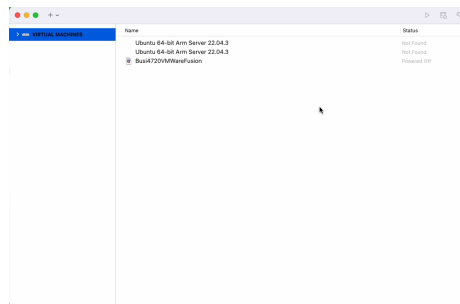
- Confirm that you wish to open the application. You may be asked to enter your password to allow this.



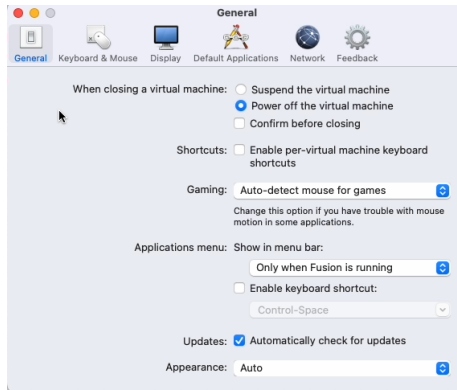
- Installation of VMWare Fusion may take a few minutes.



- When installation is complete, VMWare Fusion Pro is launched and will present you with a list of virtual machines or a welcome screen if there are none on your computer.



- Select "Settings..." from the VMWare Fusion main menu bar.

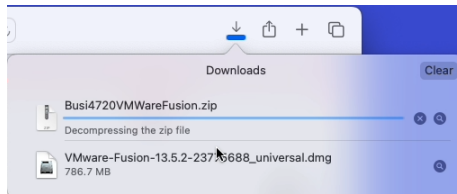


- In the Display settings, ensure that resizing the virtual machine is selected both for single window and full screen modes.

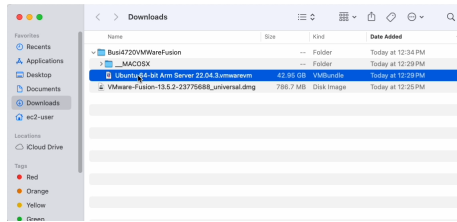


- Download the [virtual machine file](#). *Warning:* This is a 15GB compressed zip file and will take some time to download.

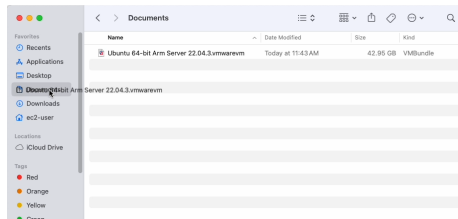
- Once the download is complete, the file will be decompressed automatically. This will also take a few minutes of time.



- Once the file is downloaded and decompressed, you will be able to see it in your Downloads folder. Expand the "Busi4720VMWareFusion" folder to see the uncompressed virtual machine VMBundle.

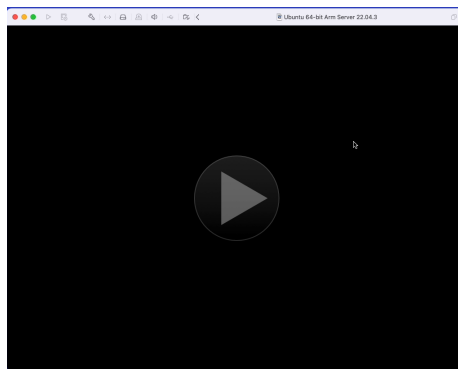


- Move this file to a permanent location, for example, to the Documents or Desktop folder.

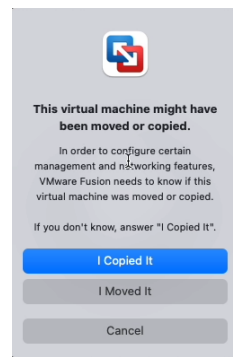


5. Start the virtual machine.

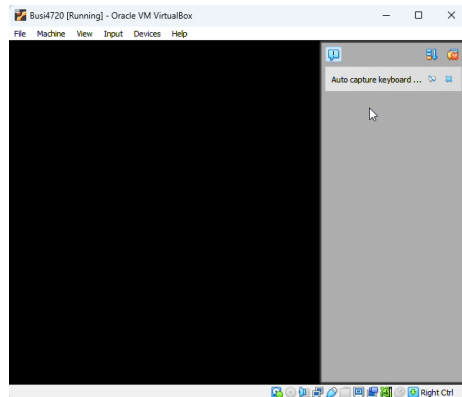
- Double-click on the virtual machine file. This will open the virtual machine in VMWare Fusion Pro.



- Press the start button to start the virtual machine. The first time you do this, you may be asked whether you copied or moved the machine. Select "I Copied It".



- Once the virtual machine has started, you will see the desktop user interface. Popular software applications are accessible via the dock on the left side of the screen. The most important application for working with this book is the Terminal window application.



- To shut down the virtual machine, select "Power Off..." from the system menu in the top right and shut down the machine.

