Concept Drift in Recommender Systems: Building a MLOps Pipeline that takes Concept Drift into account

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# Kurzfassung

Gegenstand

**Schlagwörter**: Pipeline, MLOps, Konzeptdrift, Empfehlungsdienst, Maschinelles Lernen

# Abstract

A style sheet

**Keywords:** Pipeline, MLOps, Concept Drift, Recommender System, Machine Learning

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# List of Abbreviations

ML Machine Learning

GPU Graphics Processing Unit

CPU Central Processing Unit

RS Recommender System

CD Concept Drift

TFX TensorFlow Extended

TFRS TensorFlow Recommenders

DCN Deep & Cross Network

DD Data Drift

STEM Academic disciplines of Science, Technology, Engineering, Mathematics

PoC Proof of Concept

IS Information System

SotA State of the Art

AI Artificial Intelligence

API Application Programming Interface

CF Collaborative Filtering

CB Content-Based Recommender System

TF-IDF Term Frequency-Inverse Document Frequency

MF Matrix Factorization

DL Deep Learning

DNN Deep Neural Networks

NN Neural Network

ANN Artificial Neural Network

SVD Singular Value Decomposition

FM Factorization Model

NLP Natural Language Processing

ReLU Rectified Linear Unit

QPS Queries per Second

CACE Changing Anything Changes Everything

CI/CD Continuous Integration Continuous Delivery

EDA Exploratory Data Analysis

# Introduction

## Motivation

Over the last two decades Machine Learning (ML) has become one of the fastest growing technical fields with an estimated growth of 21% in 2022 compared to its previous year (Rimol, 2021).

It managed to secure a position as one of the top fields in computer science for scientific research as well as enterprise adoption. ML combines concepts of linear algebra and statistics and applies them to large datasets to find patterns and generalizations in the data, which can be used to make predictions or classifications.

Leveraging these complex algorithms with the computational power of modern GPUs and CPUs, ML has seen application in a large variety of sectors ranging from medicine for diagnostics, to transportation for self-driving cars and e-commerce for shopping cart optimization (Choy et al., 2018). The latter sector employs so called Recommender Systems (RS) with the goal of suggesting products that coincide with the taste of the customer. With the advent of e-commerce, RSs have gained increasing interest from academia and especially the enterprise sector (Singh, Choudhury, Dey, & Pramanik, 2021). RSs serve a major role for large tech corporations in engaging, retaining and enticing the user-base of their platform (Jannach & Zanker, 2022). Netflix for example uses its own RS to suggest its users what movies they might be interested in. In order to incentivize research in the field of Recommender Systems, Netflix introduced the Netflix Prize in 2007: A dataset was made public with users and their movie ratings. The goal was to create a RS that would beat Netflix’ own RS at that time. For this challenge a prize pool of $1.000.000 was written out (Bennett, Lanning, & others, 2007). To this day datasets of movie ratings remain a popular way to benchmark RSs.

Despite the wide use and success of Recommender Systems and Machine Learning in general, it still is a relatively new field with a lot of research opportunity (Jordan & Mitchell, 2015). While Recommender Systems are considered integral to many online-platforms, their precision and accuracy often lack in comparison to other ML fields. This is, among other things, due to the nature of the data that Recommender Systems work with, which is often sparse (Khusro, Ali, & Ullah, 2016). Consequently Recommender Systems are especially susceptible to bad data quality and therefore could profit from comprehensive data curation and monitoring. This lends itself to take a data-centric approach when building, deploying and maintaining a RS, which is one of the subject matters that the field of MLOps sets out to tackle (Miranda, 2021).

MLOps emerged from the paradigm of DevOps and seeks to apply an automated and standardized approach to the lifecycle of ML applications, similar to what DevOps does for conventional Software. MLOps is attuned to the specific needs and problems of Machine Learning, such that its practices vary from those of DevOps, while still sharing the same goal of rapid and frequent deployment of Software (Makinen, Skogstrom, Laaksonen, & Mikkonen, 2021). The effect of data quality on the ML model presupposes that data quality management is an integral part of every MLOps system, since data quality affects all aspects of the machine learning lifecycle (Renggli et al., 2021). Detrimental data to the ML system’s performance can manifest itself in different ways.

One manifestation is concept drift (CD), which describes a changing outcome *y* to a constant input *x* over time (Lu et al., 2018). Deteriorating RS performance due to CD can directly impact the health of the online platform it is used on, as outputs of RSs are generally reciprocated back to the user experience. For instance, if a movie streaming platform stops recommending appropriate movies to a user because it failed to adapt to the change in taste, the user might stop watching movies on that platform and eventually cancel their subscription. Issues of Concept Drift need to be addressed and mitigated to ensure user-base retention for online services. Additionally, it needs to be incorporated into a MLOps system to benefit from the maintainability, consistency and automation of a unified process.

The product of this work, called an artifact, will be the implementation of a concept drift-aware MLOps pipeline for a RS. CD-awareness meaning, that it possesses the ability to account for potential CD in the data.

This paper serves as a thorough documentation of the design of the artifact, which is based on a comprehensive dissection of scientific literature touching the topics of Recommender Systems, MLOps and Concept Drift. The result will then be qualitatively evaluated and discussed.

This research follows the design science research (DSR) methodology of Alan R. Hevner (Hevner, March, Park, & Ram, 2004).

## Research Methodology

Design science is a research paradigm that emerged as a differentiation to natural science in STEM. Natural science, also referred to as behavioral science, is associated with fields like mathematics, physics, biology and chemistry. Its research methodology follows the objective of uncovering facts and theories about reality. Juxtaposed to the natural science lies the design science. Instead of uncovering rules and theories about the nature of reality, design science sets out to engineer and create artifacts with tools from scientific literature. Design science is predominantly represented in the engineering and computer science fields, where proof of concepts (PoC) and prototypes are the result of a lot of academic works. Both behavioral science and design science have distinguished approaches on how to conduct research.

Design science research contains a set of frameworks and best practices to manage academic work in the design science department. One of the more prominent methodologies is Alan R. Hevner’s “three cycles” of DSR (Hevner et al., 2004). Hevner originally designed his framework to involve the research aspect more closely to the development process of Information Systems (IS) in enterprise environments. It consists of 3 cycles which are closely related to each other and serve to build an artifact. The three cycles are what Hevner argues separates design science from other research paradigms (Hevner, 2007).

The artifact is the eventual product of the academic work using DSR. Since its first publication in 2004, DSR has found application in a wide variety of fields that surpasses conventional engineering and computer science. This means that the term “artifact” has a broad definition and is consequently hard to delimit. Generally, an artifact means anything that emerges from design science research. It could range from a theoretical model that was derived from other academic work, to a physical prototype or a production-ready software system.

The goal of DSR is to create an innovative artifact, which incorporates both theoretical-scientific, as well as the practical-environmental (e.g. business) aspects into its design. The iteration through the three cycles creates a mutual feedback-loop between the artifact and the science and business environment (Hevner et al., 2004). The result is an artifact, which is attuned to the business needs of an enterprise, while also holding scientific value and enriching the academic field with new insights and findings.

In the following, the three cycles will be elaborated in more detail.

1. **The relevance cycle**: The relevance cycle initiates the DSR process. In this cycle all requirements relevant to the artifact and the research are worked out. First, a problem is defined and opportunities and arguments are laid out supporting research to resolve the problem with an artifact. Since DSR has its roots in the enterprise sector, it is vital to map out and contextualize the environment this research takes place in, as it directly influences the design of the artifact. In order to evaluate the artifact, acceptance criteria need to be defined. This way a conclusion can be made whether the artifact succeeded in its goals to improve the environment or not (Hevner & Chatterjee, 2010). In this work the results of the relevance cycle are to be found in the introduction of this paper.
2. **The rigor cycle**: Following the relevance cycle, comes the rigor cycle. While the relevance cycle establishes the requirements for the project, the rigor cycle introduces the methods, drawn from scientific literature, used to create the artifact (Hevner & Chatterjee, 2010). This so called *knowledge base* consists of engineering methods and scientific theories and sets the foundation from which the artifact will be designed and built from. A thorough rigor cycle ensures that the artifact is grounded on state-of-the-art literature from the academic field. This establishes the connection to other scientific contributions and thus sets it apart from routine designs and routine design processes (Hevner & Chatterjee, 2010). The rigor cycle gives the artifact the scientific weight it requires to be acknowledged as an academic contribution, consequently it is vital that the design of the artifact draws sufficiently from the knowledge base of the rigor cycle. The knowledge base is referenced in the State of Research of this paper
3. **The design cycle**: “*The internal design cycle is the heart of any design science research project.*” (Hevner & Chatterjee, 2010) The design cycle is the culmination of the relevance and the rigor cycle. The information acquired from the two prior cycles will now be deployed to design and implement the artifact. Hevner points out that it is not possible to retain both maximum relevance and rigor simultaneously, thus a balance between both need to be struck (Hevner & Chatterjee, 2010). In the context of the design cycle the rigor represents the actual construction of the artifact, meaning the implementation of the knowledge base gained from the rigor cycle. In opposition to the rigor stands the relevance. The relevance represents all the requirements and evaluation criteria that were specified in the relevance cycle. The discrepancy between the relevance and the rigor gets resolved by the artifact, which is the bridge between both and thus constitutes the business and scientific contribution. The design cycle is documented in the Artifact design chapter of this paper.

DSR is a non-linear process. With progression of the research project, the relevance, rigor, and design cycle can change as new insights are garnered. Through iterative cycles Hevner’s DSR methodology accounts for the often unpredictable nature of the artifact creation process. Should either parts of the relevance, rigor or design fall out of line with the current state of the project, it needs to be updated by reiteration (Hevner et al., 2004).

Once the research is conducted and the artifact is created it itself becomes part of the knowledge base, whose insights can now be used for other research projects.

# Current Environment & State of Research

## Environment

Machine Learning serves great value to businesses. In 2017, Netflix for instance claimed an estimated saving of $1 billion through their use of RSs (Columbus, 2017). Since 2017, ML algorithms became more sophisticated and hardware more powerful to make Artificial Intelligence (AI[[1]](#footnote-2)) operations more efficient, effective and in return more profitable. Gartner calculated the revenue of the AI software market to be over $51 billion in the year 2021 with a prediction to surpass $60 billion by the end of 2022 (Rimol, 2021). Open-source software, ML cloud services and an active community make AI more accessible to a wide variety of businesses. These developments make a growing number of institutions consider optimizing, augmenting, or even reinventing their current operations with ML. McKinsey’s “The state of AI in 2021” reported that 56% of their surveyed businesses have adopted ML and AI in at least one of their business functions. An increase of 6% compared to the preceding year (Chui, Hall, Singla, & Sukharevsky, 2021). It’s apparent that ML and AI receive increasing interest in the enterprise sector.

According to a survey conducted by Refinitiv, out of 447 international institutions that use ML, only 46% have deployed AI in multiple areas and are core to its business, whereas 44% deployed ML in pockets, while the remaining 10% were still prototyping and investing in its infrastructure (Baker, 2019). This indicates that a majority of enterprises, while considering or using ML for their businesses, struggle to embed it into their existing infrastructure. This observation is also supported by Algorithmia’s 2020 report on enterprise machine learning, which uncovers that 55% of companies “*actively developing machine learning lifecycles or […] beginning their machine learning journey*” (Algorithmia) have yet to deploy a machine learning model. This report highlights that a lot of the main difficulties of ML lie in its operational aspect, such as reproducibility, versioning of models and scaling of the ML system. This leads to “*unreasonably long roads to deployment*” (Algorithmia) and impedes evolving the ML system to higher levels of maturity (Algorithmia). Another Refinitiv study also identified lacking data quality to remain the biggest challenge for ML and data science (Refinitiv, 2020).

The aforementioned McKinsey whitepaper made the observation that companies most successful with AI were employing advanced operation procedures such as MLOps, as well as putting greater effort into mitigating and reacting to “*AI-related-risks*”, such as concept drift. This applies to the operation of RSs as well. Operationalizing the ML lifecycle and ensuring high model quality through data quality assurance are essential to a RS in a productive environment. A data-centric approach to RSs combined with the application of MLOps practices should prevent or alleviate unsatisfactory RS performance while ensuring maintainability and reproducibility to an mostly automated ML process. Especially the phenomenon of concept drift needs to be accounted for as it is an inevitable occurrence for a majority of real world data. A key challenge for RSs is its often unpredictable nature and sudden appearance, which can be detrimental to business operation. Shift in the data can be subliminal as it is not tangible in most cases and therefore can go unnoticed by Data Scientists and ML engineers. Consequently the effects of CD can first become apparent through degradation of model performance. Scalable and automated ways to account for change in ML data still are in their infancy and therefore require further research and development to engineer solutions. In the long run a state-of-the-art ML infrastructure will make RS operations more profitable for businesses through labor reduction and performance improvements.

This introduction poses the baseline for the relevance cycle of Hevner’s design science research method. The collected insights from the whitepapers serve to map out the environment and establish the motivation for this research. The acceptance criteria for the later artifact will be in part derived from the findings and motivation in this chapter, which will be elaborated in the following chapter.

Having established the environment and motivation, the following research question is formulated for this work: “*What can a MLOps pipeline for a recommender system, that takes concept drift into account, look like?*”

## Recommender Systems

### Overview

Online platforms like Amazon, Netflix and Facebook sit upon massive amounts of data garnered from their user-base through interactions on their websites and applications. The internet enabled firms to leverage information about customers in a large-scale, centralized and automated fashion. In the past, in order to collect insights about customers, polls had to be conducted on sight of a physical store. The results were costly polling ventures of small sample sizes over a limited time period. Nowadays large quantities of high quality data is streamed uninterruptedly into data lakes where they are stored. This unprecedented amount of data enables data science and machine learning to thrive. While many large corporations live in abundance of user data, it still remains a main challenge for them to gain useful insight from the data (Cai & Zhu, 2015).

Recommender Systems use the given data to uncover preferences of end-users to then tailor the service to their likings. The most known examples are shopping basket recommendations on e-commerce sites like Amazon or movie and video recommendations of streaming platforms like Netflix and YouTube. As a generalization, the term *item* is used to denote any object in a set of items that can be recommended to a user, such as a product or a movie. The recipient of an item is referred to as the *user* or *query*. The user-item relationship makes up the foundation of any recommender system, as RSs work under the assumption that there exists dependencies between user- and item-centric activities.

The main task of a RS is to correctly identify these dependencies and use them to match appropriate items and queries (Aggarwal, 2016). In order to reduce information overload on websites with video and product catalogues, RSs are applied to identify items that will interest the user (Alyari & Jafari Navimipour, 2018). Aggarwal (Aggarwal, 2016) lists four aspects that define a good recommendation:

1. *Relevance*: The most fundamental objective of a RS is to recommend items that are relevant and interesting to the individual user.
2. *Novelty*: The RS should introduce items to the users that they weren’t aware of before. It has been shown that recommender systems can negatively impact the sales diversity of e-commerce, when only popular items are recommended to users (Fleder & Hosanagar, 2007).
3. *Serendipity*: Similar to the point above, RSs should have the ability to surprise users with somewhat unexpected but relevant items. While novelty focuses on the quantitative aspect of recommending items that aren’t on the radar of the user, serendipity considers the qualitative-psychological impact of a surprise in the recommendation.
4. *Increasing recommendation diversity*: This last point envisions that RSs should ensure a diverse set of items in a recommendation feed. This should be done to not fatigue the user with items of similar categories. An example would be the recommendation of movies from a variety of different genres.

Generally speaking, Recommender Systems

Throughout the years, different approaches have been developed to optimize recommendations and overcome obstacles in this field. These recommendation systems are broadly categorized in content-based RS (CB), collaborative filtering (CF), knowledge-based RS and hybrid RS (Aggarwal, 2016). In the following, CF and CB are introduced to provide an overview of the RS landscape, as both belong to the most popular research topics for RSs, excluding hybrid systems, which are a combination of multiple stand-alone RS methods. Collaborative filtering will be elaborated in more detail, as it is a key concept for the RS implemented in the artifact. Figure 1 shows the structure and an overview of the recommenders based on Aggarwal (Aggarwal, 2016). Highlighted in red is the type of recommender system that will be used in this this research project, which is a deep and cross network. Before elaborating CF and CB, there will be a chapter dedicated to retrieval and ranking, two tasks RSs are used for. This is chapter is then followed by a chapter about RS data.



Figure 1: structure and overview of RSs derived from Aggarwal (2016)

### Retrieval & Ranking

Most RS algorithms are employed to fulfill either a retrieval or a ranking task. The retrieval task deals with the generation of appropriate candidates for a recommendation. Within a dataset, a subset of similar items and users are identified (Fernández-Tobias, Cantador, Kaminskas, & Ricci, 2012). The objective is to narrow down the dataset to the relevant users and items, that are then used by a ranking algorithm to make scoring predictions. Opposed to the retrieval task, the ranking task makes concrete recommendations to a user. Ranking can manifest itself in form of a predictive score, or a top-items list (Google LLC, 2020b, 2020c).

Neighborhood algorithms and other similarity measures are popular methods used for retrieval operations, whereas state-of-the-art ranking is often done via matrix factorization or deep learning models. Neighborhood algorithms, like k-nearest-neighbors, can be used for ranking as well, as explained in chapter Collaborative filtering.

In practice, both retrieval and ranking are often combined into a joint RS, as seen in Figure 2. Since ranking algorithms are computationally more expensive, they wouldn’t scale well in an enterprise setting. For this, the faster retrieval operation pre-selects a set of candidates, which are then run against the ranking model.

Retrieval systems are outside the scope of this research project and thus won’t be discussed in greater detail. All references about RS in this paper are related to ranking models.

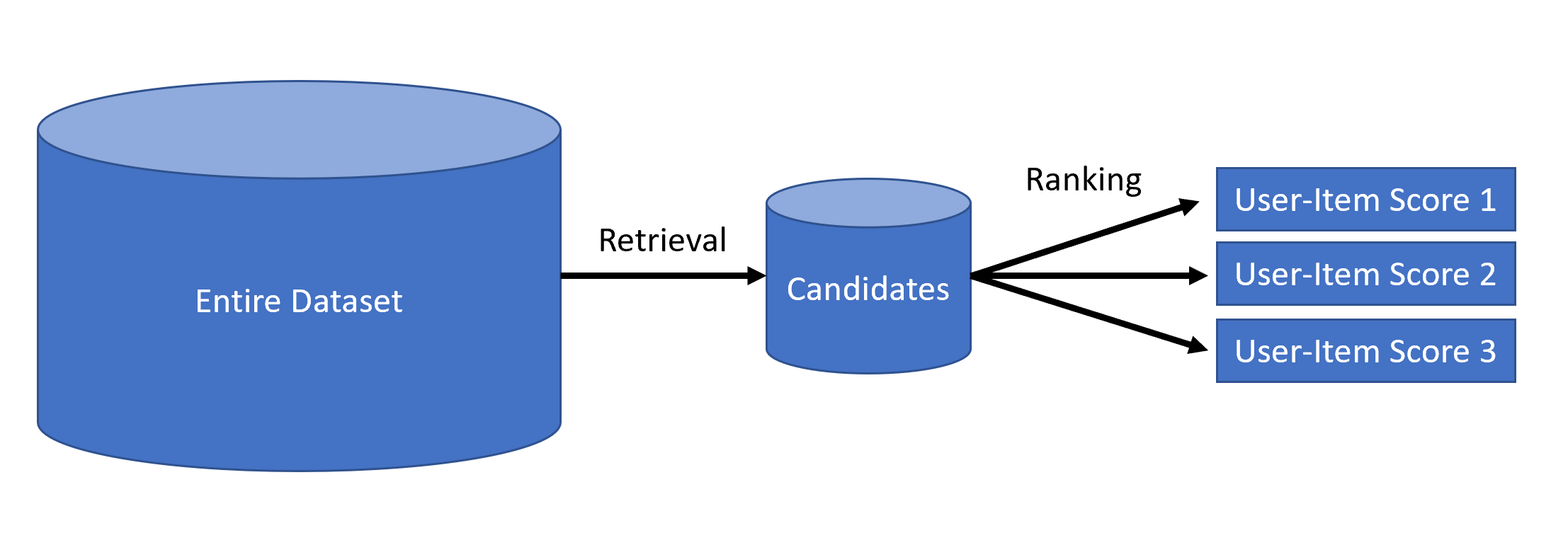


Figure 2: Example of retrieval and ranking in a recommender system

### Data

The data used by RSs often differ from those of other Machine Learning tasks, as it is often sparse. Sparsity refers to very few user-item interactions relative to the size of the whole dataset. Assuming one would construct a matrix of an e-commerce website, wherein represents the entirety of its user-base and represents all the items in the product catalogue. Every time a customer buys or inspects a product, an interaction between the user and the product is denoted in . Since customers only interact with a fraction of the offered products in an e-commerce web shop, there are very few entries in the 2-dimensional matrix, hence it is sparse. This is a fundamental problem that RSs face and try to overcome.

The value of an interaction between item and user can be represented in different ways depending on the context of the recommendation. Is an interaction defined as a review with a rating between 1 and 5 stars, it will be marked with an integer value in a range of 1 to 5. These rating types allow the user to quantifiably express their like or dislike of an item. These are called interval-based ratings (Aggarwal, 2016). Another popular rating type is the unary rating system. These are used on social media platforms like Instagram and Twitter. These ratings are encoded in a binary fashion, wherein an interaction (e.g. like) will be conventionally denoted with the value 1 and abstinence will be denoted by the value 0 (Aggarwal, 2016).

Both the like-feature as well as the 5-star rating system are called explicit ratings, as they are actively and consciously given by a user. While explicit ratings generally provide reliable information about an users opinion, it remains a challenge to entice users to rate. This in return makes the data more susceptible to shilling attacks, where item ratings get skewed by coordinated bombardments of overly positive or negative ratings by external sources (Khusro et al., 2016).

In opposition to explicit ratings, lie the implicit ratings. These rating systems derive a rating from the user through their behavior. An examples could be the watch time of a movie. This example works under the assumption that people that don’t close a movie, are engaged with it. The drawback of an implicit rating is its ambiguity, as online behavior needs to be interpreted first and then translated into a rating. User behavior can be interpreted wrongly and skew the data. Continuing the aforementioned movie example, a user who slept during the movie out of boredom, would be misinterpreted as a user who was engaged throughout the whole movie, since they didn’t close it. An implicit rating system has the advantage that it is not reliant on the user to give feedback, which generally results in more ratings. This is a significant factor considering sparsity is a main challenge for RSs (Su & Khoshgoftaar, 2009).

In a recommender dataset the value of a user-item interaction is referred to as a *sparse feature*. They can be augmented with so-called *dense features*. Whereas sparse features only occur through user-item interaction (e.g. a rating), dense features are specific attributes that describe the item or user at hand, comparable to features in other ML tasks.

Data understanding plays a vital part in the success of recommender systems. There are many variables ranging from its rating system over its features and sparsity that determine, which recommender system is the most suitable for the given task (Aggarwal, 2016, p. 128).

### Content-based recommender systems

Content-based methods make recommendations based on a user's past interactions with items. These users are referred to as objects and contain information from their history of item-interactions (e.g. movie watches, ratings, etc.) (Lü et al., 2012). These items have descriptive attributes, named content, which form the basis of which these recommender systems derive their recommendation decisions from (Aggarwal, 2016). These contents are used to picture the preferences of a user. In a movie recommendation setting such a content could be the genre of a movie. Depending on what contents are found in an object, certain assumptions can be made about a user's likings. Other items can then be assigned to the user based on a similarity measure between the object and other item objects. Alternatively new items can be suggested by weighing the content by their frequency in an object. Popular methods for content-based filtering are information retrieval algorithms like TF-IDF (term frequency-inverse document frequency) (Lü et al., 2012). One characteristic of CB is that both the item and the user are represented in the same feature space, in order to measure similarities between an user object and all item objects.

Item recommendations can be done independently from other users, as it is only of interest how the individual user is positioned to the items in the vector space, which is one of the advantages it has over CF. This means that CB can be applied in environments with relatively small user-bases. Since CB relies on a user history to make recommendations, it has challenges finding the right recommendations for new users, since there isn’t enough content to map out the user (Aggarwal, 2016). Conversely, making recommendations for new items is relatively easy, since the item objects only need to be provided with appropriate content. The success of content-based filtering is highly determined by the quality of attributes in these objects. These features need to be cautiously engineered. Since the predictions are calculated on the basis of the user’s and item’s attributes, it is not possible for these algorithms to uncover hidden variables in the data, like for instance some CF algorithms could. In general, the performance of CBs are very reliant on how the data has been engineered by data and ML engineers.

### Collaborative filtering

CF has been the baseline for RSs for years. In September 2009, the BellKor team was awarded with the grand Netflix prize, which it has won by applying a CF model on the competition dataset (Koren, 2009a). As the name implies, CF uses the "collaborative" power of the user-item interactions in the dataset (Aggarwal, 2016). Instead of solely relying on item and user descriptions, CF uses the composition of the dataset to determine the output of a recommendation. Colloquially, one would say that the recommendation is driven by a "joint effort" of every user-item interaction. Unlike CB, recommendations can be made without relying on hand-engineered features (Google LLC, 2021).

**Memory-based filtering.** CF can be divided into *memory-based* methods and *model-based* methods. The prior uses neighborhood based algorithms to make recommendations. These methods belong to the earliest CF practices, where ratings are predicted based on similar users or items (Aggarwal, 2016). "Memory-based" relates to the fact, that the dataset is loaded into memory, in order to make calculations. The dataset is then actively used to compute the right recommendation through nearest neighbor methods. For that reason the data in memory-based CF can be described as part of the model, as recommendations are determined by the arrangement of datapoints within the vector space.

Memory-based methods can either take a *user-based* or *item-based* approach. In user-based CF, rating predictions are provided by like-minded users. Similar user groups are identified using nearest neighbor techniques, then the unobserved ratings are filled in by the values of the peer group. This could be done by calculating a weighted average of all ratings from that group (Aggarwal, 2016). In item-based CF a set of items a user has rated most similar to an unrated item are retrieved with a neighborhood algorithm. In item-based CF a set of items, already rated by a user, are used to determine the value of an unspecified item. Via neighborhood algorithms items are determined that resemble the unrated item the closest. Like with user-based CF, the rating is derived from the neighboring entities, only that instead of using the values from similar users, the values from similar items are chosen (Aggarwal, 2016).

The advantage of a memory-based method is its interpretability, as ratings are directly taken from other users or items. Since CF methods (memory-based and model-based) obtain their values through other users and items, it is easier to achieve serendipitous recommendations, as similar users might have interacted with items the target user isn’t familiar with.

Memory-based algorithms don’t work well with sparse data, as there might not be sufficient items or users similar to each other (Aggarwal, 2016).

**Model-based filtering.** Unlike memory-based approaches, model-based methods have a training phase, whereby a model is learned. The goal is to create a function (the model) that approximates the results of an item-user interaction, which is conventionally achieved through optimization of a loss function with gradient descent or alternating least square algorithms (Koren, Bell, & Volinsky, 2009). Two of the most prominent model-based CF techniques are matrix factorization (MF) and deep learning (DL).

The former has been the standard in the field of RS and was also utilized by the winning team of the Netflix prize in 2009 (Koren, 2009a). MF is foremost a dimensionality reduction technique with which the user-item matrix gets decomposed into *low-rank[[2]](#footnote-3) latent factors*. These latent factors are the components that make up the *latent factor model* (LFM), which is a low dimensional representation of the initial matrix. LFM, broadly classified as a *factorization machine* (FM), works under the assumption that there are underlying latent variables in the data. Latent variables are variables that are not directly observable but can be inferred through mathematical computation, like for example singular value decomposition (SVD) (Loehlin & Beaujean, 2017). Koren et al. (Koren, Yehuda and Rendle, Steffen and Bell, Robert, 2022) state that the observed rating values are due to effects associated with either user or items, independently of their joint interaction. Therefore, there are large item and user biases embedded in the data. An example would be the tendency for some users to give either higher or lower ratings on average, or conversely products that receive systematically higher or lower ratings. These “hidden” factors can be retrieved during model training and incorporated into a latent factor. In the training process the latent factors are distilled that best depict these propensities in the data (Vellido, Lisboa, & Meehan, 2000). In LFM, the rating of an unobserved user-item interaction is constructed by applying the dot product of factor of user and factor of item (Aggarwal, 2016). While the baseline LFMs only use the sparse matrix of user-item interactions, more sophisticated variants, like SVD++, can also incorporate dense features and implicit ratings into its model, which allows it to factorize more information and further improve prediction quality (Koren et al., 2009).

The main advantage of MF lies in its ability to algorithmically detect latent vectors in the dataset. Overarching correlations are extracted in the dataset and used to make predictions. Unlike CB methods, the models don’t rely on hand-engineered features to make recommendations, which evades the human-error component of feature engineering and speeds up the development process. Dot products are computationally light, which makes LFM models scalable and applicable on large datasets (Blondel, Ishihata, Fujino, & Ueda, 2016). The performance advantage of FMs rapidly diminish when modeling higher-order feature combinations however, which is a tradeoff to allow for more complex embeddings in the model (Blondel, Fujino, Ueda, & Ishihata, 2016). FMs being bound by their shallow structure hinders their overall representative power (Wang, Fu, Fu, & Wang, 2017). In practice, simple FMs still are very capable at making accurate predictions (Dacrema, Boglio, Cremonesi, & Jannach, 2021).

In order to circumvent the above-mentioned limitations of FMs, *deep neural networks* (DNN) have gained increasing interest in RS research over the past years (Dacrema et al., 2021). *Neural networks* (NN) are very effective at retrieving complex embeddings from their input, which makes them state of the art in the field of image recognition and natural language processing (NLP).

Neural networks emulate a simplified model of how human or animal neurons actually work in a brain. Like the brain in biology, *artificial neural networks[[3]](#footnote-4)* (ANN) consist of multiple individual neurons that are interconnected, whose joint response to an external input creates an output. Unlike neurons in our brains, which are structured in a complex and organic way, neural networks are arranged in layers, which contain a set of artificial neurons (Charniak, 2019).

The baseline neuron in NNs can be thought of as an individual mathematical function, that consists of a weight component and a bias component, which are wrapped inside an activation function. This neural function receives an input, which can either be the external input data, or the output of neurons from the previous layer, depending on whether the neuron sits in the first layer of the network or not. In the neuron, the input value gets multiplied by the weight then the bias offsets the product by a certain value. The activation function then refactors this weighted sum, which is the output value of the neuron. Activation functions are non-linear functions that determine how “activated” a neuron is, i.e. how large the output value should be. A popular activation function is *rectified linear unit* (ReLU), which represses any activations (i.e. neuron output value of 0) up to a certain threshold (Nair & Hinton, 2010). In the output layer (the last layer of the network) activation functions like *Softmax*[[4]](#footnote-5) are preferred, as they normalize the neural computation to a probability distribution. In the case of classification, an *Argmax* function can then be applied on the output layer to retrieve the neuron with the highest Softmax output, which represent the class the NN predicts with the highest confidence (GOODFELLOW, BENGIO, & COURVILLE, 2016). The non-linearity of these activation functions in combination with the multiplicity of artificial neurons, allow NNs to retrieve highly non-linear patterns from the data, which is the reason they are also referred to as universal function approximators (Charniak, 2019; Gurney, 2014; Hanin, 2019; Heaton, 2012).

NNs are learned through *deep learning*. Intuitively speaking, the objective of the learning process is to tweak each parameter (i.e. weight and bias) in a NN in such a way that a desired output corresponding to a certain input is generated. The training is done through an iterative process called *gradient descent*. Before the training process begins, each weight in the network gets assigned an initial random value. This offsets the NN to learn distinct embeddings, which would not be possible if each weight received the same value. During the training process *test data* is fed through the network. With a loss function the desired target output is compared to the actual output of the unoptimized model . The goal with each training iteration is to decrease the distance between and . To achieve this, the gradient vector of the loss function is calculated with respect to the entire NN. This computation of the gradient in a NN is what’s called *backpropagation*. We established earlier, that each neuron (excluding the first layer) is a function of the outputs from the neurons in the previous layer. Backpropagation recursively calculates the gradient inside each neuron in the NN, starting with the output layer and propagating through the entire network to the input layer. Having calculated the gradient, the opposite sign of it is taken and multiplied by the *step size*, a coefficient which defines how aggressively the parameter values are tweaked. This process is repeated until the NN function converges to a local minimum. A small step size increases the learning time, since it takes longer to reach a minimum. Conversely, a large step-size runs the risk of exceeding the local minimum, which results in inferior performance (Gurney, 2014; Heaton, 2012; LeCun et al., 1989).

For every model training, it is crucial to separate the training data from the test data to prevent an overfit of the model. This is especially true for NNs, as they have the tendency to perfectly adapt to the data it gets trained with, which greatly impacts the generalizability of the model and impairs prediction performance on unseen data (May, Maier, & Dandy, 2010). In general, the learning of NNs is a non-trivial task, as they are very sensible to the embedding of the input data, the initialization of the weights, the choice of activation functions and hyperparameter tuning, like step size and *dropout regularization[[5]](#footnote-6)*. A lot of variables that determine the efficacy of a model make it harder to extract the full potential out of a NN. In addition, NNs can’t easily be interpreted because of their complexity, which is why they are often treated as black boxes, especially with very deep models. (Dacrema et al., 2021) discovered that simple MF methods show similar performance to other state of the art DL models, when selecting the right parameters. Compared to a linear LFM, NNs also have a lower processing speed, which is detrimental to large recommendation datasets, since passing a neural network is much more computationally expensive than a simple dot product calculation (Rendle, Krichene, Zhang, & Anderson, 2020).

This notion changes when comparing NNs to polynomial FM, which are much more computationally complex compared to their low-degree counterparts (Blondel, Fujino, et al., 2016). As universal function approximators, DNNs have the potential to extract much more complex patterns and relationships out of the dataset, especially as both dense and sparse features can be trivially incorporated into its embedding. With ongoing research, DNN approaches for RSs are further being refined and its deficiencies, like performance, investigated and improved. Currently, high profile enterprises like Google use NNs in large scale environments for recommendation tasks (Covington, Adams, & Sargin, 2016).

### Deep & Cross Networks

When using data with multiple features to make recommendations, *feature crosses* become an important concept. The notion of feature crosses is that the combination of multiple features into one feature can yield an expressive predictor variable, that would otherwise not be extractable using only single features. Individual features that have very low correlation with the label can therefore reveal themselves to be highly correlative in conjunction with other features. Uncovering these *cross features* allows to extract additional information out of the data, which in return can significantly improve model performance. The amount of features that are embedded in a cross feature is denoted by the order: A cross feature being the product of two features is called a cross feature of 2nd order. Cross features of 3rd order and up are referred to as higher-order features (Wang et al., 2017; Wang et al., 2021).

Suppose a RS that predicts the user rating of a movie. Besides the sparse features (in this case the user ID and movie ID), the data also contains the genre and length of the movie as dense features. Assuming that length as a stand-alone feature is not a strong predictor for the rating of a movie, it could be combined with the genre feature into a separate cross feature of 2nd order. Now, a feature that has both variables encoded in it can be used to better predict the ratings of a movie, the hypothetical interpretation being, that different genres set certain expectations of what the movie length should be. For instance, while it might be acceptable or even desired for fantasy movies to be 3 hours long, a 3 hour runtime for a comedy could generally be viewed unfavorably.

Feature crossings are often done as part the data preparation phase of a ML lifecycle (Studer et al., 2021). This process can be very daunting and time consuming, as it entails trial and error of which cross features are best suited for accurate predictions. Also, feature crosses aren’t always obvious and thus might go unnoticed. Instead of manually feature engineering cross interactions, it would be lucrative to directly incorporate feature crosses into the ML model and let the learning algorithm determine the weights of cross features. DNNs are capable of learning arbitrarily high order functions, provided the NN is deep enough. They however only learn feature interactions implicitly, which means that they do not reliably pick up on higher feature crosses without drastically increasing the network size (Wang et al., 2017). This brings the general efficiency of DNNs into question, especially in light of large scale recommendations that need to compute high amounts of queries per second (QPS) (Rendle et al., 2020).

In order to reduce the overall model size of NNs and invoke explicit cross feature learning, Google proposed their deep and cross network (DCN) in 2017 (Wang et al., 2017). In addition to a classic feed-forward multilayer perceptron (MLP) NN, as covered previously, a DCN is augmented by an additional cross network, consisting of cross layers. The cross network enables learning on bounded-degree cross features, whose order is explicitly defined by the layer depth of the cross network. This way, ML engineers can now directly incorporate the order of feature interactions they wish to embed, by changing the cross network structure. At the start of the network (i.e. the input layer) the feature vector doesn't have any explicit cross feature interactions yet and is therefore of order 1. With each cross layer the feature vector passes, the maximum polynomial degree of cross interactions increases by 1. This is because for each cross layer the output of the previous layer gets crossed again with the input vector to construct a new cross term . A visualization of a cross layer in Google’s DCN V2 architecture is seen in Figure 2, whereby the learned parameters and respectively denote the weight matrix and the bias of the cross layer.



Figure 3: Visualization of a cross layer (Wang et al., 2021)

A DCN is always made up of two components: A cross network and a conventional MLP neural network. While the cross network is used to learn explicit bounded-degree feature interactions, the DNN component is used to learn implicit feature interactions in the data[[6]](#footnote-7). With both components the model is therefore able to effectively leverage explicit and implicit interactions in the data, which improves prediction performance. Depending on what architecture is chosen for the DCN, the DNN can either follow the cross network, or run in parallel, as seen in Figure 3.



Figure 4: Possible DCN architectures (Wang et al., 2021)

With the introduction of cross layers, the model becomes more effective and predictable at embedding feature interactions into its network. While feature selection and feature embedding still remains a non-trivial task, it is only of concern at the level of individual features, since cross feature extraction is now delegated to the model training, instead of being a separate, manual task (Shan et al., 2016). This saves labor time and removes the human error element. Being able to extract both implicit and explicit feature interactions in the data through the combination of a cross network and DNN, makes DCN overall more competitive at predicting ratings. Besides improving the prediction performance of recommendations, the model also becomes smaller and in return faster, which makes it more viable for large-scale RS operations.

### State of the Art Technology

**TensorFlow Recommenders** (TFRS) is a python library built on the Keras API. TFRS unifies and simplifies the building of DL RS models under TensorFlow, by providing functionality and components at each step of the RS development cycle, such as methods for retrieving and ranking recommendations, activations functions, and ready-to-use RS datasets (TensorFlow, n.d.). TFRS also provides built-in RS specific network layers for model building, such as the aforementioned cross layers. In combination with the accessibility of the Keras API, it opens the possibility to intuitively realize state-of-the-art DCNs (TensorFlow, 2022a, 2022b).

**TorchRec**. In January 2022, Meta released an open source library for building recommendation systems under PyTorch, called TorchRec. This library was utilized to build the state-of-the-art RSs employed in Facebook’s Newsfeed and Instagram’s Reels (Meta, 2022). TorchRec provides optimized RS kernels for efficient matrix multiplication in the model. CUDA enabled GPU acceleration improves model training time, provided that the supported hardware is used (Zhang et al., 2022). TorchRec can automatically split up large RS models into shards for distributed prediction serving. *Sharding* is a common pain point for large scale RS operations. Modern RSs can’t be hosted on a single server, as the model size exceeds most memory specs of server racks. In order to retain the performance of large RS models, the model need to be broken up into shards. These shards are then distributed to multiple compute nodes, where they can jointly compute and serve predictions (Lui et al., 2020). Meta seeks to address this topic by providing automated sharding plans for models (Greenberg, Taylor, Ivchenko, Liu, & Sudarshan, 2022).

## MLOps

### Problem

Machine Learning has its origins in academic research (Minsky, 1961; Rosenblatt, 1961). Over the last twenty years it also found its way into the enterprise sector, where it now stands as the backbone of the most successful companies and most profitable business ventures. With the transition from academia to industry, ML engineers and data scientists are confronted with unique sets of challenges that are usually absent in a sterile research environment. While research often is dedicated to investigate one aspect of ML (i.e. the algorithm), enterprise devotes itself to the development, operation, and maintenance of entire ML systems. Software in research is written to investigate a certain topic and to accept or decline a very specific hypothesis. It has a very narrow use-case and therefore lacks generalization, which makes it often not suitable for a productive setting. In addition, development of the software usually stops with the conclusion of the research project (Garousi, Petersen, & Ozkan, 2016). On the other hand, software employed in business environments are built for longevity, with their goal being a profit driver for the company. This is composed of low operational costs and high revenue-yield (Heemstra, 1992; Myers, Kappelman, & Prybutok, 1997). As a result of these different goals, both research and industry employ different development paradigms, with focus on different aspects respectively. These differences are observed in the RS landscape of academia and industry, for example. While in academia state-of-the-art RS algorithms are assessed with their performance on static benchmark datasets (e.g. MovieLens dataset), industry also considers the integration of the algorithm into the system architecture. Instead of merely relying on benchmark performance, industry systems need to take into account the scalability of an algorithm and embed their RS into their application in a meaningful manner (Covington et al., 2016).

One of the first papers to illustrate challenges of productive ML systems is Sculley et al. (Sculley et al., 2015), which uses the framework of *technical debt*, found in software engineering, to describe problems in a productive software environment. To be more precise, the term technical debt refers to ongoing maintenance cost due to shortcuts taken during initial development of the software. The debt analogy is used to visualize upfront cost-savings due to cheaper and faster development, that however need to be paid back with interest down the line with excessive software service and maintenance (Cunningham, 1993). In classical software engineering technical debt can manifest itself in different ways, such as code refactor, deleting outdated modules, reducing dependencies and improving documentation (Fowler, 2019). The same phenomenon of technical debt is also observed in ML systems. Sculley et al. (Sculley et al., 2015) points out that a lot of the technical debt incurred in ML, is due to the fact that it directly interacts with the external world. Since ML holds a tight bond with the ever-changing data it consumes in real-world applications, maintenance turns out to be particularly difficult and expensive. So, in addition to the established technical debt of software engineering, ML also receives its own set of technical debt that needs to be considered as well (Sculley et al., 2015).

Enterprise ML are often very complex software systems that comprise various interconnected components, consequently the ML algorithm itself only makes up a small portion of the entire system. In addition to the complex array of variables in the ML model alone, all parameters from the other software components have an effect on the ML system as well, which results in a web of entanglements. It becomes hard to keep track of all dependencies within the system, which, in return, complicates the maintenance of the software, which is referred to as *configuration debt*. The changing anything changes everything (CACE) principle goes into effect and any adjustments to the system can severely compromise the model performance (Sculley et al., 2015). Related to the configuration debt, is the *reproducibility debt* in ML systems. Reproducibility is an inherently difficult undertaking in productive ML systems, as these algorithms often work with randomized parameters, non-determinism in parallel learning and continuously changing data.

Another form of technical debt occurs when ML researchers and data scientists don't adjust their habits when moving from the academic field to industry. As mentioned in the beginning, academia and industry apply different software development paradigms. Not adapting their ways of working to the needs of production software may greatly impact the maintainability of a ML system. Common culprits are so-called *glue code* and *pipeline jungles*. Glue code is understood as software code, that doesn’t directly contribute to the functionality of the software’s or system’s requirements, but acts as a compatibility layer between two or more components. Glue code is often written in presence of stand-alone ML packages or commercial-of-the-shelf software products, in order to integrate them into the ML infrastructure. Glue code generally distracts from the actual purpose of the application. It makes code less readable and degrades performance. It also makes the software less flexible to changes and updates, as glue code is tailored towards specific interactions between specific components (Sculley et al., 2015). The pipeline jungle is a special type of glue code, which evolves out of a base pipeline, through incremental expansion. Often found in the data pre-processing phase, these pipelines are continuously built upon to accommodate for new data sources and data transformation steps. In the end, the pipeline becomes error prone, hard to test, and convoluted (Sculley et al., 2015). Both glue code, and pipeline jungles are symptoms of code that isn’t designed to be reused and built upon, which is emblematic of software in research projects, as they often serve as PoCs.

Lastly, a large portion of the technical debt in ML systems is due to their nature of working with dynamic data from the external world. Not dealing with changes that occur in the dataset can lead to an amalgamation of technical debt, to the point where it might even render the whole model obsolete (Lu et al., 2018; Sculley et al., 2015). The implications of changing data will be dissected in greater detail in chapter Concept Drift.

### What is MLOps

Paying down ML related technical debt demands a re-examination of status-quo practices in a business environment. Industry has different sets of priorities compared to academia, which need to be respected. Instead of determining the viability of a model based on specific performance metrics alone, the effects on operability of the whole system need to be taken into consideration. Small performance gains at the cost of maintainability usually don't fare well in an enterprise environment. At the same time, a infrastructure needs to be set in place, that is capable of preventing and mitigating technical debt. This entails sustainable design patterns in ML code, such as modularity and abstraction, rigorous testing, as well as practices that promote team work (Sculley et al., 2015). The goal for the last few years, was to apply practices of agile software development to ML. As assessed in the prior chapter, ML development contains a few peculiarities, that sets it apart from conventional software development, which makes it incompatible with common technical methodologies, like DevOps. Out of necessity to employ methodologies that are attuned to the unique characteristics of ML, the term MLOps (was born. Despite gaining attention from high profile ML researchers and companies, MLOps still remains a vague term that is hard to pin down (Kreuzberger, Kühl, & Hirschl, 2022; Tamburri, 2020). This is due to the novelty of this field, with different sources defining MLOps differently.

A Google whitepaper about this topic defines MLOps as “*a methodology for ML engineering that unifies ML system development (the ML element) with ML system operations (the Ops element)*” (Salama, Kazmierczak, & Schut, 2021, p. 5). Similar to DevOps, MLOps seeks to employ practices and tools to shorten development cycles and enable rapid iterations of ML software (Symeonidis, Nerantzis, Kazakis, & Papakostas, 2022). Streamlined operational and governance processes are used to increase performance, reliability and security of ML systems. To sustain this agile development, focus is put on collaboration methods and tools. One overarching theme of MLOps is the unification of research, which is inherent to ML, with the efficiency of modern software development to make ML more profitable and scalable in real world applications (Salama et al., 2021; Sculley et al., 2015).

Tamburri (Tamburri, 2020) defines MLOps from a technological perspective as a set of middleware and software components, that ought to be orchestrated in the cloud to realize at minimum 5 specific functions:

* Data ingestion & transport
* Data transformation
* Continuous ML (re-)training
* Continuous ML (re-)deployment
* Output production & presentation to the end-user

These points are derived from popular cloud MLOps platforms, like Kubeflow, which enable scalable ML pipeline orchestration through containerization (Kubeflow, n.d.). These platforms provide a large portion of the infrastructure to realize state-of-the-art productive ML systems, that can be broken down into following areas of responsibility:

**Automation**. Due to changing data that ML systems consume, they need to be constantly updated and deployed. Employing a CI/CD (continuous integration/continuous delivery) approach for ML, allows development teams to quickly release updated iterations of models and other components. In CI/CD, new models can be automatically compiled into a build and be deployed instantaneously in the needed environment.

CI/CD is often realized in conjunction with automated pipelines, which are a vital part of successful ML operations. A typical ML lifecycle usually consists of repeating steps. Taking the CRISP-ML process as an example, every development cycle needs to run through a data preparation phase (Studer et al., 2021). Since these steps remains consistent throughout different lifecycles, they ought to be automated. An automated process is usually represented by a DAG (directed acyclic graph), which models a process with nodes and edges. The DAG is read by a workflow management tools, such as Apache Airflow, which executes specific tasks in a defined order (Kreuzberger et al., 2022). Pipelines remove manual labor from the ML process, which speeds up the development of new models, reduces the human error element and makes the process more consistent.

In state-of-the-art MLOps the conventional CI/CD practice is expanded by *continuous training* (CT). CT aims to automate the entire model training process. The ML system should be able to detect deteriorating model performance and initiate a model retraining pipeline (Denis Baylor et al., 2019). One form of achieving CT is deploying entire ML pipelines instead of a trained model. This way, instead of deploying a single ML model at the time, an entire pipeline can be deployed to recurrently create new and optimized models (Denis Baylor et al., 2019).

**Reproducibility**. The most straight-forward implementation of reproducibility is through versioning of the source code with a *version control system* (VCS), like Git. As an industry standard, VCSs allow to trace and revert changes in the code. This way previous states of the software code can be restored and rebuilt (Serban, van der Blom, Hoos, & Visser, 2020).

For ML applications, code versioning alone is not sufficient however. Since model training is not only dependent on the model configuration, but also the training data, the dataset needs to be versioned and tracked as well, in order to replicate a previous model (Ruf, Madan, Reich, & Ould-Abdeslam, 2021). This can be achieved through time travel functionality of modern data lakehouse architectures, such as Databricks delta lake (Yavuz & Chockalingam, 2019). Alternatively there are dedicated data management tools, like Rok from Arrikto, integrated into Kubeflow (Arrikto, n.d.).

As alluded to earlier, ML utilizes a wide array of hyperparameters and randomization-seeds for learning. These need to be tracked as well, in order to ensure reproducibility of the model. These are stored in a meta data store, alongside other information, such as training duration and training date.

**Monitoring**. Continuous monitoring is used in MLOps to assess the health of deployed ML systems. Most importantly, the prediction quality of served models are frequently measured in an automated fashion. Important metrics are given a threshold value, that are tracked live by the monitoring system. Sophisticated monitoring implementations automatically trigger pipelines to take corrective actions, upon noticing insufficient model performance (e.g. a re-training pipeline). The observations made during monitoring can feed back into the consecutive model to make it more robust. Automated monitoring can be supported by human monitoring through dashboard visualizations on applications like Kibana, Grafana, or PowerBI (Kreuzberger et al., 2022).

### Maturity Levels

Maturity levels determine the sophistication of a MLOps implementation in an productive environment. The maturity of a ML System can be defined by its technological infrastructure, employed methods of operation, and automation through pipeline orchestration. Because of the novelty of the field of MLOps, there isn't one universal maturity model agreed upon by the scientific community. Companies that are heavily invested in AI therefore design and develop their own maturity models. The most prolific proposals stem from Microsoft and Google (Symeonidis et al., 2022).

**Microsoft**. Microsoft’s maturity model is divided into 5 levels. Level 0: No MLOps, Level 1: DevOps but no MLOps, Level 2: Automated Training, Level 3: Automated Model Deployment, Level 4: Full MLOps Automated Retraining (Figure 4). In this model the following factors determine the maturity of the ML system: People, model creation, deployment, and application integration (Microsoft, n.d.).

Level 0 represents the baseline maturity, where no MLOps is employed. Systems of this maturity have little to no automation in the ML lifecycle. Builds of the ML application are created and deployed manually. Because of missing model repositories, reproducibility is limited. ML models are often treated as black boxes, that are developed in an enclosed environment by an isolated team. Consequently, the ML system isn’t embedded into the larger infrastructure of the company and it is therefore hard to retrieve feedback post deployment.

Level 1 maturity ML systems incorporate standard DevOps practices into the ML lifecycle. Opposed to level 0 maturity, builds are created and deployed automatically. There exists a baseline testing infrastructure for unit and integration tests of the application. Production models are now tracked through a basic model repository, though experimentation still is done in an ad-hoc fashion. Data engineering receives more attention and is brought together with the data science team. Data is collected continuously at this stage. The ML team is however still separated from the software engineering team. For this reason the ML model still remains a black box.

While level 1 maturity focuses on the automation of software engineering tasks relating to the application, level 2 also puts focus on automating the data science tasks. The data engineers and data scientists work together to create unified pipelines for data preprocessing and model training. Scoring scripts and experimental code are now tracked to ensure reproducibility of models. Models are still manually released to the software engineering team for application integration.

Level 3 maturity foresees unification of ML and application. Software engineers work with data engineers to automate model integration into the application. Opposed to lower maturity levels, models are now automatically released to the software engineering team through CI/CD pipelines. Every step from data collection to deployment is traceable. All ML and application components are tested automatically. Deployed models are evaluated with testing methods such as A/B testing.

Maturity level 4 envisions a mostly automated ML lifecycle. ML development and application development are fused into one process. Software engineers integrate live monitoring of various performance metrics into the application. This way model performance can continuously be checked in the production environment. Based on these live metrics a retraining pipeline can automatically be initiated, known as CT.

**Google**. Google’s maturity model is divided into 3 levels. Level 0: Manual process, Level 1: ML pipeline automation, and Level 3: CI/CD pipeline automation (Figure 4). Like in Microsoft’s model, a higher level means a more sophisticated implementation of MLOps for the ML system (Google LLC, 2020a).

Similar to Microsoft’s maturity model, level 0 denotes the baseline maturity, where no MLOps practices are employed. Every step is done manually, usually executed interactively on a notebook. There is no distinction between experimental and productive code, as iterations of the model are done on the same notebook until a servable model is trained. No CI/CD is employed, which means that testing is usually done in an ad-hoc manner within the notebook. Once the model is deployed there isn’t a monitoring infrastructure to observe the model performance in production. Consequently, level 0 ML systems don’t deploy models regularly and run the risk of not noticing degrading prediction quality.

The goal of level 1 ML systems is to perform continuous training. In level 0 a trained model is deployed. In level 1 a complete ML pipeline is deployed, which triggers a model training process. The pipeline then continuously delivers up-to-date models that are then used by the application. In production, the input data is monitored live to detect potential shifts and skews, which then can trigger the pipeline to retrain a model. In order to improve reusability and accelerate the pipeline development process, the source code is broken up into modular components. There is now a strict separation between development and production environment.

In Level 2 ML systems, CI/CD is employed to the full extent. The test and deployment of pipelines are automated and the experimentation process is orchestrated. Intelligent monitoring is used on live data, which is able to discern whether a model retraining or a new experiment cycle should be initiated. Information about models, data and metadata are stored in a model registry, feature store[[7]](#footnote-8) and metadata store. This makes ML pipelines reproducible and transparent.



Figure 5: Levels of Microsoft's (left) and Google's (right) maturity model (Google LLC, 2020a; Microsoft, n.d.)

### Concept Drift

“*One of the things that makes ML systems so fascinating is that they often interact directly with the external world. Experience has shown that the external world is rarely stable. This background rate of change creates ongoing maintenance cost.*” (Sculley et al., 2015)

One manifestation of ongoing change in data is called *concept drift* (CD). Specifically, CD describes a changing outcome to a constant input over time (Ng, Crowe, & Moroney, 2021). Real world examples of CD could be changing house prices due to a fluctuating house market, or people changing their taste in movies because of aging or genre trends. These changes in the underlying distribution of ML data are often unforeseeable. Therefore they are hard to detect and often go unnoticed for a long time. CD runs the risk of first being noticed through deterioration of model performance in a productive environment.

(Gama, Žliobaitė, Bifet, Pechenizkiy, & Bouchachia, 2014) identify 4 common types of CD as seen in Figure 5:

**Sudden drift**. The distribution[[8]](#footnote-9) of the data undergoes an abrupt change in a short amount of time that persists indefinitely. This phenomenon is also known as *concept shift*. One example for concept shift would be the drastic drop in demand for commercial flight to a country where war just broke out (International Air Transport Association, 2022).

**Gradual drift**. This type of concept drift has two active concepts in the data. Over time, the probability of the initial distribution decreases and gets gradually displaced by the new distribution. An example could be the genre representation in the billboard top 100 charts. As new genre trends emerge, old ones fade out and get replaced by the new ones. During the transitory period both concepts coexist in the data.

**Incremental drift**. Unlike gradual drift, the incremental drift can’t reliably be ascribed to a discrete amount of distributions. The drift occurs slowly with a lot of variables and concepts at play. Consequently, incremental drift is only noticeable when looked at over a long time period. Cultural and societal changes are often incremental in nature.

**Reoccurring concepts**. Previously seen distributions might reappear after some time, as an irregular occurrence. While seasonal changes are reoccurring, they wouldn’t necessarily be classified as concept drift, as they are predictable to an extent. An example for reoccurring concepts are rare weather phenomena like hurricanes (Knotek & Pereira, 2011).

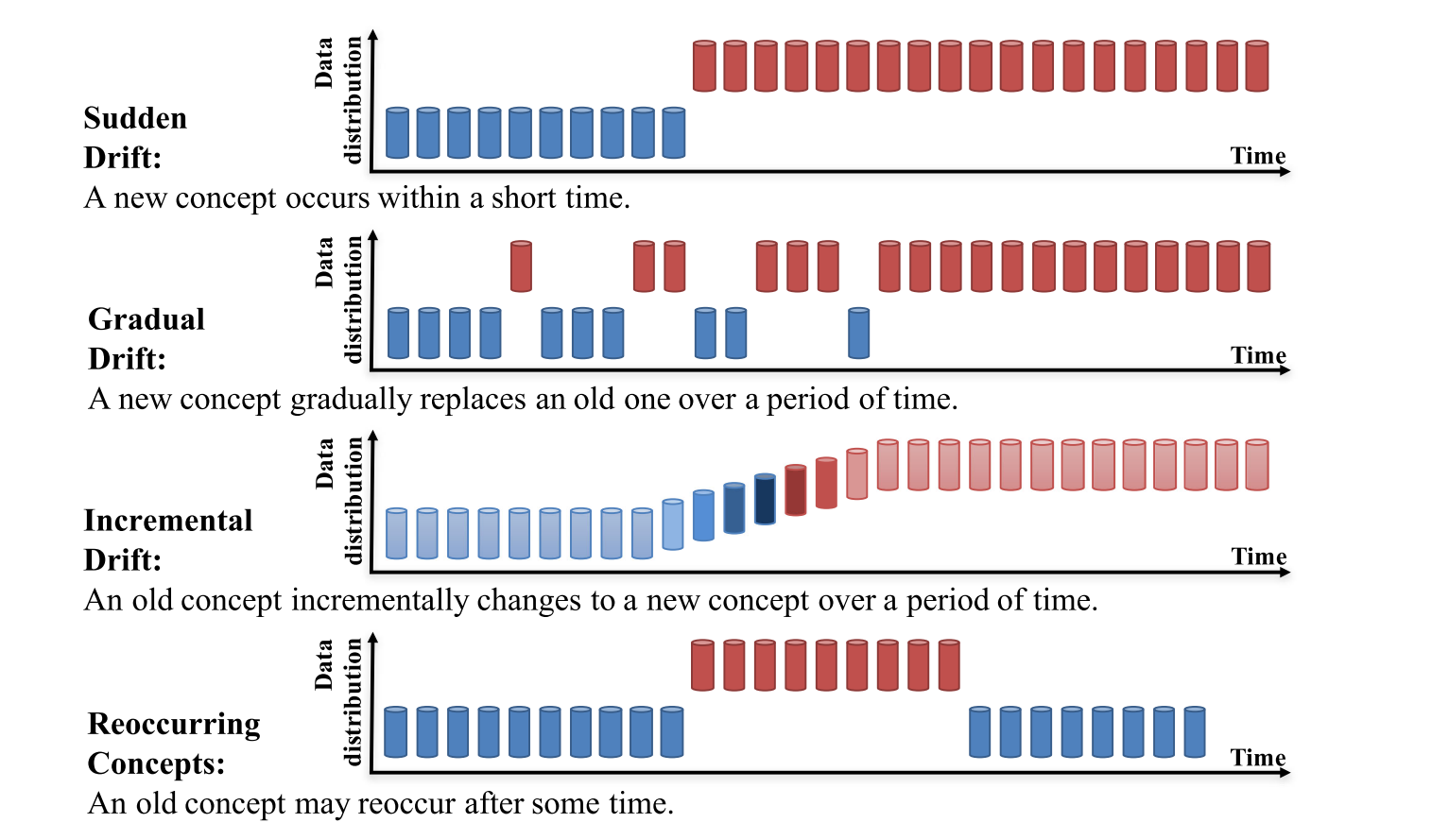


Figure 6: Types of concept drift (Lu et al., 2018)

Not all changes in data can be attributed to CD. A major challenge in CD detection is to differentiate true CD from noise and outliers. Noise and outliers are random deviations in the data, that aren’t indicative of any larger patterns and shouldn’t therefore be treated as CD (Gama et al., 2014).

Throughout the years, researchers have developed frameworks and algorithms to deal with CD. (Lu et al., 2018) proposes a framework to integrate CD handling into a production ML system (Figure 6). In this proposal CD handling consists of 3 parts: CD detection, CD understanding, and CD adaptation.

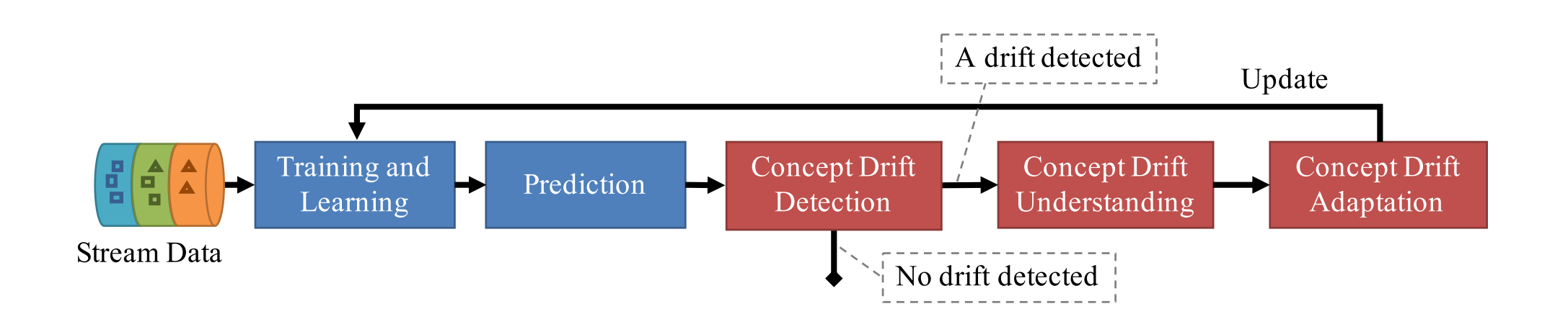


Figure 7: Framework for handling concept drift in machine learning (Lu et al., 2018)

**CD detection**. CD’s unpredictable and inconspicuous nature makes it often hard to identify. CD detection employs techniques and mechanisms to characterize and quantify CD. CD can be uncovered in different ways.

The most common way to detect CD is through observation of model performance in the productive environment. This type of CD detection is referred to as error *rate-based drift detection*. A performance decrease over time is indicative that characteristics of the data have shifted. Different algorithms can be employed to analyze CD using the model performance metrics. Drift Detection Method (DDM) was the first algorithm to assess the severity of the CD at hand, based on how the model performed over a set time window (Gama, Medas, Castillo, & Rodrigues, 2004). The window defines the timeframe of past model performance, which the current model performance is compared against. If DDM detects a significant error-rate increase, compared to the old data, it’ll declare that CD has occurred. Built upon DMM, other algorithms have been developed, such as the Early Drift Detection Method (EDDM) and the Fuzzy Windowing Drift Detection Method (FW-DDM) (Baena-Garcıa et al., 2006; Liu, Zhang, & Lu, 2017). In the context of automated CT pipelines, it becomes especially important to configure appropriate values for CD detection. The right performance metrics need to be monitored and the right performance thresholds need to be selected. Additionally, the performance needs to be observed over the right time window. Are these values not set correctly, CD detection might become too sensitive to noise. Conversely, the CD detection might pick up on CD too late.

Instead of using model performance to determine CD, the dataset can be analyzed. This category of algorithms are called *data distribution-based drift detection*. In this case, severity is quantified using distance functions to measure dissimilarities between new and old data (Lu et al., 2018). Density based algorithms and Principle Components Analysis (PCA)-based algorithms can be employed for CD detection (Feng Gu, Zhang, Jie Lu, & Chin-Teng Lin, 2016; Qahtan, Alharbi, Wang, & Zhang, 2015).

**CD understanding**. CD understanding deals with the extraction of important information about the CD at hand. CD understanding usually is the output of CD detection algorithms. Additional understanding can be acquired by employing additional methods, like visualization. According to (Lu et al., 2018) following questions should be answered in order to gain a comprehensive CD understanding:

* When did CD occur and how long does it last? (When)
* How severe is the concept drift? (How severe)
* What were the drift regions of CD? (Where)

Identifying when CD occurs is the baseline requirement for every CD detection method and therefore the most trivial question to answer. Detecting CD implicitly contains information about when the CD at hand was detected. Through continuous monitoring of the data and model, the duration of CD can be measured as well.

In order to satisfy the second question, CD needs to be quantified in a non-binary fashion. The severity of CD is measured by most dedicated CD detection algorithms, like DDM. As addressed in CD detection, the severity can be derived from diverging model performance and dissimilarities between old and new data. The steeper the performance drop, or the bigger the distance between the data old and new data, the higher the severity level of the CD.

The most difficult to answer question is *Where*. Drift regions are regions of conflict between the new and old distribution. Feature vectors located in a drift region are affected by the CD and contribute to the error-rate of the model. Identifying drift regions makes CD more tangible and allows for a more in-depth analysis of model performance. The techniques to make out drift regions are however highly dependent on the employed drift detection model. This means that there is no generalizable way to map out drift regions in the feature space (Lu et al., 2018).

Data understanding doesn’t need to be collected algorithmically alone. Insights into occurring shifts in the data can be visualized as well (Koren, 2009b). Visualizations can analyzed by data scientists to qualitatively assess CD in a dataset.

**CD adaptation**. The most conventional way of reacting to CD is to trigger a model retraining process. In the context of MLOps this is referred to as CT. After the defined performance threshold is not met, a new model is trained with new data. There are variations to simple retraining. One is called *ensemble retraining*. After CD is detected a new model gets trained, which is then added to an ensemble with the old model. Both models are then used to make predictions. Ensemble retraining is suitable for gradual CD, where two concepts are present in the data.

CD adaptation can also be incorporated into the model itself. Gradual forgetting is a technique employed during model training, where training examples are weighted based on how new they are. The result is a model that is more sensitive to the characteristics of newer data (Koychev, 2004). Another way of incorporating CD adaptation is to manually embed temporal effect into the ML model (Lo, Liao, Chang, & Lee, 2018). (Koren, 2009b) identifies temporal dynamics throughout the whole dataset and incorporates them into a CF model. This research was conducted on a Netflix dataset, where it discovered that movies receive better ratings as they age. This observation can be converted into a mathematical formula and be incorporated into the RS model.

### State of the Art Technology

In order to overcome the challenges of MLOps, a wide variety of tools have been developed in the past years. Particularly, two notable open source end-to-end ML platforms are used in industry to operationalize large scale ML applications: *TensorFlow Extended* (TFX) and *MLflow*.

**TFX**. TFX is maintained by Google and used for their own ML projects. TFX compartmentalizes common tasks of a ML lifecycle into separate components. The entire ML process from data ingestion, data evaluation, data transformation, model training, model evaluation, up to model deployment is standardized and modular. Currently there are a total of 10 built-in components that are provided by TFX (Figure 8). The repository of base components can also be expanded by building custom components.

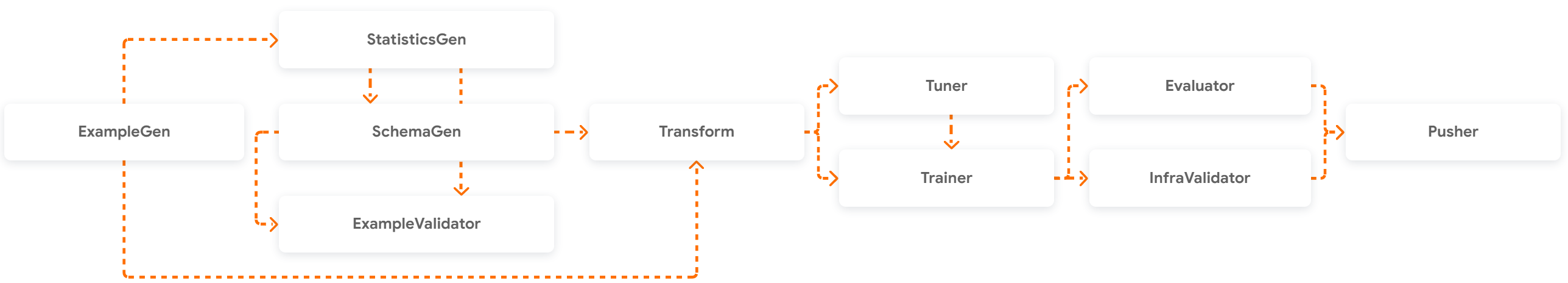


Figure 8: TFX components (Google LLC, 2019a)

These components can be orchestrated and run in a pipeline with support for Airflow and Beam. In addition to the components TFX includes libraries and software components for metadata storage, model creation and model evaluation (Baylor et al., 2017). TFX’s central metadata storage is called ML metadata. This SQLite database acts as a model repository and feature store. In addition, it tracks training parameters and the order of component execution to make every pipeline run traceable. ML metadata also saves progress of model training, in order to continue the training at a later time (Crowe, 2019).

Custom components gives TFX the flexibility to embed external libraries into the pipeline, however TFX is clearly optimized to work within the TensorFlow ecosystem. This means that the TF API, TF Model Analysis (TFMA), TensorBoard and TFRS are natively supported by TFX and work without extensive configuration.

The following will provide an overview to the essential components of TFX (Google LLC, 2019b):

*ExampleGen* is the first component of the TFX pipeline and ingests data in a specified directory. It supports various data types, ranging from TensorFlow’s TFRecord to CSV and popular Big Data file formats like Avro or Parquet. The ingested data is then versioned and stored as TFRecord.

*StatisticsGen* takes the saved data from the ExampleGen component and generates rudimentary statistics from it. This component serves to provide a general overview of the data, which can also be used by consecutive components.

*SchemaGen* automatically retrieves and infers the schema of the dataset at hand.

*ExampleValidator* takes the outputs from StatisticsGen and SchemaGen and searches the dataset for anomalies that might impact training performance. This component investigates the data for missing values, data skews between training and serving data, and potential data drift.

*Transform* is used for data preprocessing tasks. Within this component, arbitrary data transformation operations can be written in a *preprocessing\_fn* function. Transform takes in the data output from ExampleGen and the data schema from SchemaGen.

*Trainer* takes on the model training process of a ML lifecycle. The Trainer component takes in a ML model file with a *run\_fn* function, which initializes the training process. For the training and evaluation data either the output from ExampleGen or Transform can be used. Training parameters (e.g. training steps) can be given as arguments. The output of this component is the trained model.

The training process can optionally be augmented with the *Tuner* component, which employs the KerasTuner module to find the best hyperparameters of a model. The output of this component is given to the Trainer component.

*Evaluator* leverages the TensorFlow Model Analysis (TFMA) library to perform model evaluation. On the basis of the automated model evaluation it can be determined whether the trained model yields superior results to the current production model. If the model supersedes the previous model it receives a “blessing”, meaning that it can be deployed to a production environment.

*Pusher* deploys the model to a specified target (i.e. directory). It takes in the model trained by the Trainer component. If the Evaluator component is used, the Pusher can determine whether to push a model based on the blessing.

**MLflow**. MLflow is another ML platform provided by Databricks. Similar to TFX, it provides a simple API to streamline the ML lifecycle. MLflow is composed of 5 main components (Alla & Adari, 2021; Databricks, 2018; Zaharia et al., 2018).

The *MLflow Tracking* component is a logging tool to track parameters, code version and metrics during any ML process. Custom log functions can be configured and ingested into MLflow Tracking.

*MLflow project* is a format to package ML source code in a modular way. These conventions standardize the code of an ML project and make it usable in automated processes.

*MLflow Models* is a format for packaging ML models to make them compatible with external tools. This component can be used to integrate models into an application.

*MLflow Model Registry* is a centralized model repository similar to TFX. It provides additional metadata on model lineage and model version.

*MLflow Model Serving* hosts models directly from the model registry, which are callable via REST API. This feature is exclusive to the Databricks cloud service.

**KubeFlow**. KubeFlow is a solution for large scale ML operations. It provides an infrastructure for a complete MLOps environment in the cloud. It comes with essential software, like notebooks for development, TensorFlow for model creation and Apache Airflow for pipeline orchestration. In addition to built-in software packages, it also provides support for various ML tools, like TFX. KubeFlow is built on Kubernetes which handles containerization and cluster computing. This cloud infrastructure is highly scalable in nature. This means that model training can be distributed between multiple compute nodes, which accelerates the training process (Kubeflow, n.d.).

**AutoML**. Automated ML (AutoML) are services that provide end-to-end ML systems. They are provided as software-as-a-service (SaaS) on a cloud platform (e.g. GCP and AWS). The internal workings of AutoML are usually hidden to the user and the system can only be configured through pre-defined interfaces. The goal of AutoML is to provide easy-to-use and intuitive ML solutions, that don’t require extensive programming skills from the user (Karmaker et al., 2022).

# Goal & Specification

## Artifact

The goal of the DSR methodology is the design and creation of an artifact. This particular work sets out to create an automated machine learning pipeline for a RS, the main research focus being the implementation of CD-awareness into the pipeline. For evaluation purposes, a specification sheet will be generated on which the artifact will be measured against. The artifact will be compared with the features listed in the specification. Based on the specifications, the final product will be analyzed, hurdles during the design and development will be identified, and the success of this research will be determined. The specifications are structured in *base specifications* and *research specifications*. While both types of requirements are integral to the whole research project, the research requirements have a direct connection to the research question at hand. During the DSR process both requirement types receive equal prioritization as both requirement types make up the whole artifact. A table with all specifications can be seen in *table 1*.

The base requirement list consists of creating a state-of-the-art RS. During this phase an appropriate dataset will be selected that the RS can be trained and run on. The RS the gets integrated into a basic MLOps pipeline without explicit CD-awareness. Throughout the iterative development process this pipeline will be expanded on to fulfill the research objective of this work.

Table 1: Artifact specification table

|  |  |  |
| --- | --- | --- |
| **Specification type** | **Specification** | **Description** |
| Base specification | Recommender system | The artifact has a SotA RS and a suitable dataset to run on. |
|  | Base pipeline | The artifact has a pipeline to run a basic ML lifecycle on. This base pipeline serves as a baseline and will be iterated over. |
| Research specification | CD detection | The artifact has the ability to detect CD. |
|  | CD understanding | The artifact has the ability to extract information out of the CD. |
|  | CD adaptation | The artifact has the ability to react to CD. |
|  | Integrated pipeline | The final artifact is one pipeline that can be executed in one go |
|  | Automated pipeline | The pipeline can run without human intervention |

The research specifications have three criteria related to concept drift, which are taken from the 2018 paper “Learning under Concept Drift: A Review” (Lu et al., 2018), presented in chapter “Concept Drift”:

*Concept drift detection* specifies the technical ability for the artifact to register concept drift in the data. An example would be the implementation of a concept drift detection algorithm.

*Concept drift understanding* specifies the ability of the artifact to retrieve additional information from the concept drift. This criteria serves to give concept drift more tangibility and helps data scientists identify the source of concept drift. The most rudimentary implementation of this specification would be a timestamp for every occurring CD. A timestamp for CD is also the minimum requirement to realize an automated pipeline, as in order to initiate concept drift adaptation the system needs to know when CD occurs.

*Concept drift adaptation* specifies an action that accounts for concept drift, like starting a retraining of the model.

Each of these components amount to what we define in this paper as CD-awareness. All three specifications will be explained in further detail in the literature review.

Besides the concept drift specifications, there are also pipeline conditions in the research specifications that envision the artifact to be one unified and automated process.

These specifications serve as an aide during the design and development of the artifact. Rather than evaluating the artifact based on how many specifications it technically fulfilled, the whole artifact will be qualitatively evaluated using the specification sheet as a guideline. In the scope of this work, one solution to CD will be implemented that is derived from the literature. It is not a comparison and evaluation of various implementations of CD-awareness. This work does not entail a quantitative evaluation of the artifact, as the focus lies on a qualitative analysis of the prototype.

## Procedure

Throughout this project various tools are used to track, organize and document this work. As part of this research, a GitHub repository was created, which contains both the thesis Word document and the software artifact.[[9]](#footnote-10) GitHub and Git are used for version control and enable work on different systems. It also serves to make the development of this research transparent and traceable. Internally, a GitHub Project Kanban board is used for project management. There, the project is broken down into individual tasks, all of which have their progress tracked. This way it is possible to gain an overview of the current status of this project and plan future development steps.

The overarching process of the artifact creation is structured in underlying phases, as seen in Figure 9. Each phase will be developed in a separate branch and then merged into the main branch, as is convention in software development.

The initial phase of this project is dedicated to the set-up of the artifact environment. This phase encompasses every necessary step to build a software environment on which the artifact will run on. The individual tasks would consist of choosing the python version, setting up a work directory and installing needed packages and other software.

After setting up the environment, an appropriate public dataset is selected for this project. If needed, a small data preparation task will be done to make the data ingestible by the MLOps tools used in this project.

Next, a base MLOps pipeline with a placeholder RS is built. It should fulfill the basic functionality of a MLOps pipeline and serves as the foundation for further development steps.

The following phase involves swapping the placeholder RS for one state-of-the-art RS discussed in chapter 2.2.

After implementing the new RS, the base pipeline then gets incrementally expanded by additional components. Each pipeline component that gets added will be tested for compatibility. Depending on what components work and don’t work, design changes to the artifact need to be made.

Having implemented the RS and the pipeline, an intermediate evaluation of the current pipeline will be done. The objective in this phase is to find out the capabilities of this pipeline in relation to the research question. What components can be leveraged to fulfill the objective of this work? Accomplishing these phases should fulfill the base specifications detailed in the previous chapter.

The following phase consists of incorporating CD-awareness into the pipeline. With the knowledge base from scientific papers and information about the environment from the whitepapers, a solution to CD will be engineered. This process will run iteratively and in accordance with the DSR methodology. Each iteration of this phase concludes with an evaluation, which will determine whether another iteration is initiated.

Once the artifact reaches a state where no additional iterations are done, a final evaluation of the artifact will be made, which concludes this research project.

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Figure 9: procedure of artifact development

## Challenges

This work is not done in association with a company, despite being a study directed towards businesses exploring a potential solution to CD in an MLOps scenario. Consequently this research can't be conducted in an existing enterprise infrastructure. This entails the challenge that, first, an environment needs to be built approximating a productive business setting. For instance, instead of using real world data from a data lake or data warehouse, this research needs to revert to publicly available datasets. Simulating a productive MLOps environment means that there is a discrepancy between this work's environment and an organization's environment, which might impact applicability of the artifact.

The open-ended nature of this work’s research question is another source of challenge that is inherent to DSR projects. In Hevner’s DSR methodology it is only specified what problem the final artifact should solve, not how it should look like, as it is part of the process to come up with an innovative design. This room for interpretation of what the artifact should look like makes it more difficult to quantify it in the end. For this reason a specification has been created, which should make the idea of the artifact more concrete. Even with a specification sheet, the challenge remains on how to meaningfully evaluate a ML pipeline. Unlike ML algorithms, there is no reference pipeline to measure the artifact against, assuming that using quantitative metrics like processing time would make sense for this work in the first place. Another aspect that hinders quantifiability of such an artifact is the phenomena of CD itself. As mentioned in the environment chapter, CD is a burden for businesses because it lacks tangibility and is therefore hard notice, let alone isolate and quantitatively measure. This project needs to deal with these limitations and find a way to evaluate the artifact in a value adding manner.

Another hurdle is a lack of scientific studies and official documentation on this specific subject, as the operationalization of ML systems only received mainstream attention in the last few years with the advent of terms like AutoML and MLOps, or tools like TFX. In the following, TFX and TFRS are chosen to illustrate the problem. This example is however applicable to other tools as well.

TFX had its first release in May 2019 (Google LLC), whereas TFRS had its initial release in fall of 2020 (Google LLC). Only in end of March 2022 has Google released its first TFX documentation using TFRS (Google LLC, 2022). As of conducting this project, more fundamental documentation is planned for both tools (Wei, 2022). It would be defensible to say that both tools still are in their infancy, going by their recent releases, their ever expanding documentation and active development. Besides the novelty of these fields, a lot of research and development in ML systems goes unnoticed by the academic community, as they are conducted behind closed doors and used as proprietary software. In 2019, Naumov, et al. stated the following about the current environment of RS research: “*Although recommendation and personalization systems still drive much practical success of deep learning within industry today, these networks continue to receive little attention in the academic community*” (Naumov et al., 2019)

# Design & Development

## Environment

### Hardware & Software environment

This research project is conducted on a deep learning compute cluster provided by the Media University Stuttgart (Theodoridis & Grießhaber, n.d.). The hardware resources are allocated by Slurm, a cluster management software. This way, multiple users can access powerful hardware to train large DL models. The DL cluster consists of multiple individual servers, which can be classified as either a CPU partition or GPU partition. While CPU compute nodes are sufficient for development and data preparation tasks, GPU acceleration can be leverages to rapidly compute tensor calculations.

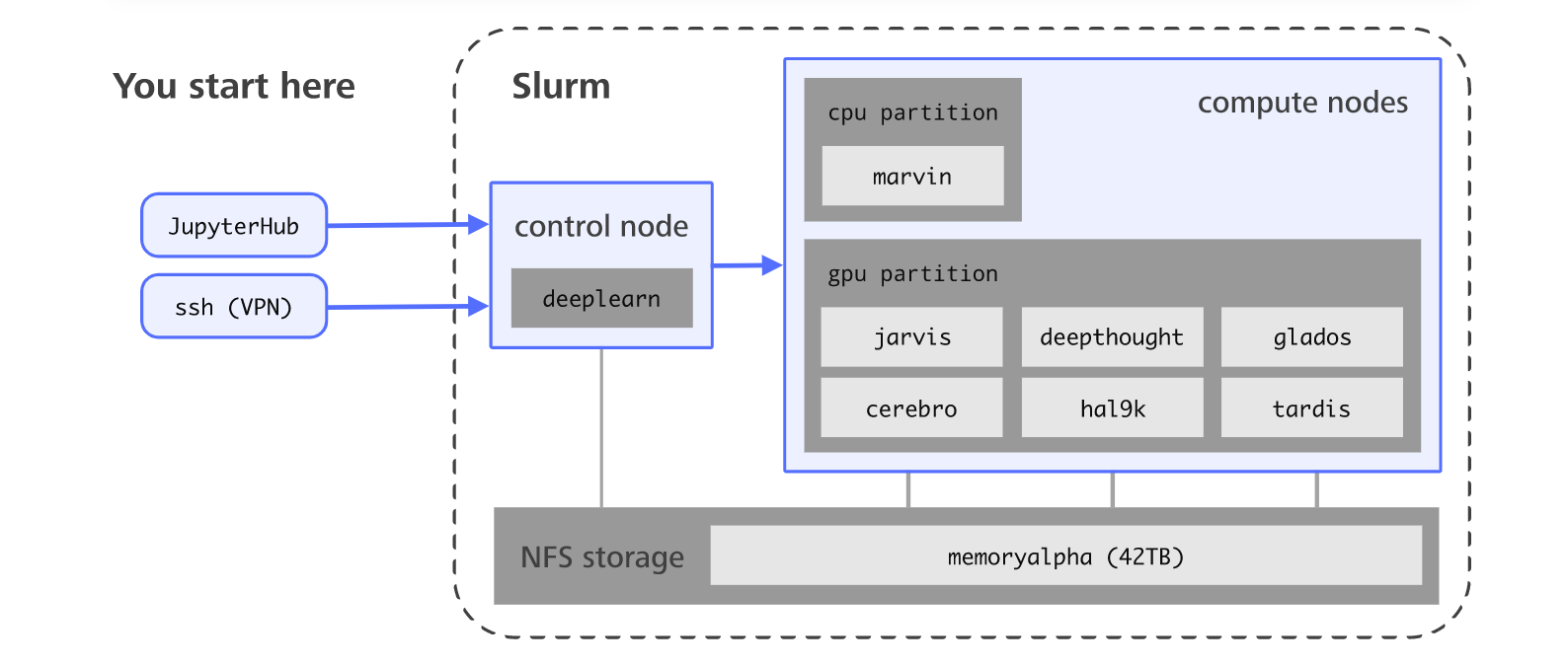


Figure 10: Infrastructure of the HdM deeplearning cluster (Theodoridis & Grießhaber, n.d.)

These servers are accessed via the control node, which acts as the entry point to the cluster. The control node can either be addressed via a web browser through JupyterHub or with secure shell (SSH), in combination with a VPN. The cluster has a shared network file system (NFS), where the files from all users are stored (Figure 9). In order to start a server, the user needs to select either a CPU or GPU instance (Figure 10). Depending on the selection, the user gets assigned a corresponding cluster node. Each time a server instance is started, an Ubuntu image gets mounted to the users work directory. This image comes with Anaconda preinstalled, in order to create and manage Python environments.

Python will be used as the main programming language of this project, as it is built-in into Anaconda and supports a wide array of data science packages (e.g. TensorFlow, NumPy, pandas) to conduct this research. Specifically, Python version 3.8 is chosen.

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Figure 11: selection of instance in the HdM deeplearning cluster

During the assessment of a suitable research environment, KubeFlow was briefly considered. As mentioned in the “State of the Art Technology” chapter in “MLOps”, KubeFlow would provide an all-in-one environment for this research. KubeFlow is a viable solution for large scale MLOps in an enterprise setting, but wouldn’t be appropriate for the prototype that is developed in this research. The set up and configuration of the environment would exceed the scope of this project. KubeFlow instantiates various containers with ML software, which are intelligently distributed across a server cluster with Kubernetes. While this makes for a scalable productive system, it also has high hardware requirements.

For this artifact, a lean software stack of few selected tools are sufficient to conduct this research.

### Working Directory

The working directory of the artifact is located in *bachelor\_2022/artifact* (Figure 12). Within the working directory are other subdirectories. The green folders seen in Figure 12 are not tracked as they do not contain any source code.

*Data* is a subdirectory where all the datasets before a pipeline are stored.

*Data\_fetch* contains scripts for data retrieval and data preparation. These scripts can be executed to download files and prepare them to make them readable by the employed software tools. The contents inside *data* are the product of these scripts.

*Pipeline\_scripts* contain all files related to pipeline execution and pipeline orchestration.

The source code for the ML models are saved in *model\_source*. The pipelines use these files to train the ML model.

*Pipeline* is the directory where all pipeline outputs are then stored.

*Sandbox* serves as an environment where concept and test code is saved. These scripts aren’t part of the artifact and thus are located in a separate folder.

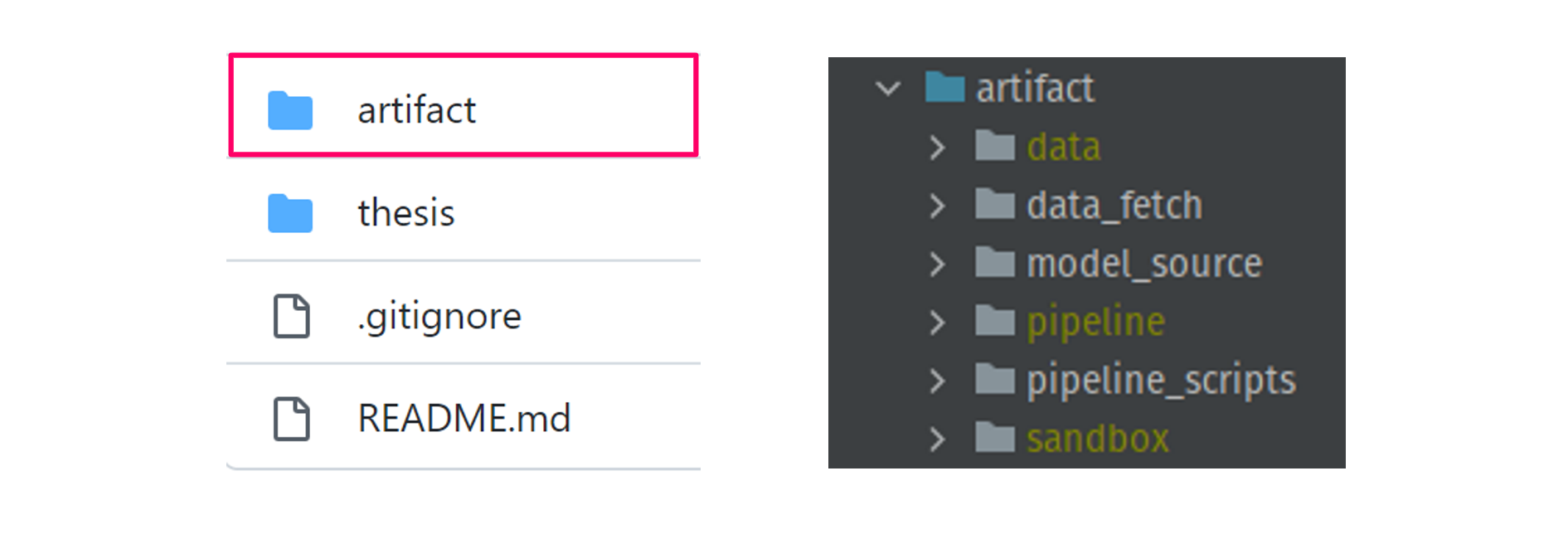


Figure 12: Working directory and its subdirectories

In the following, the tool selection and the implementation of the individual artifact components are discussed in greater detail.

## Data

### Dataset Selection

This research is not conducted within a company environment, which means that a public dataset needs to be chosen in order to realize the artifact. There are various public datasets available that are used for benchmarking by the scientific community. One of the most popular datasets for RSs are the MovieLens datasets from GroupLens (Harper & Konstan, 2016). GroupLens collects movie ratings of users since July 1998, that are then continuously released to the public. They offer datasets of varying sizes ranging from 100.000 up to 25 million ratings, wherein each user has at least 20 movie ratings. An overview of the different datasets can be seen in Table 2.

Typical for a recommender dataset, MovieLens entail information about user and item interactions. The two smallest datasets, 100k and 1m, have additional demographic information about their users. These dense features make them a viable choice in light of SotA RSs that can leverage both sparse and dense features for predictions.

This project sets out to create a PoC MLOps pipeline. This means that RS benchmarking isn't part of this research. For this reason, larger datasets, like 20m and 25m, have not been considered, as the training process would have taken up a significant amount of time. Due to the time constraints of this project, it is important to be able to make rapid iterations to the pipeline. We determined that for this reason, the MovieLens 100k dataset would be sufficient for this type of research.

Table 2: Overview of MovieLens datasets (GroupLens, n.d.)

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Name** | **100k** | **1m** | **latest-small** | **20m** | **25m** |
| **Ratings** | 100.000 | 1.000.209 | 100.836 | 20.000.263 | 25.000.095 |
| **Users/ Movies** | 943/ 1.682 | 6.040/ 3.952 | 610/ 9.742 | 138.493/ 27.278 | 162.541/ 62.423 |
| **Dense user features** | Yes | Yes | No | No | No |
| **Timespan** | 1997-1998 | 2000-2003 | 1996-2018 | 1995-2015 | 1995-2019 |

### Dataset Description

The MovieLens 100k dataset can be divided into 3 parts. The first part maps the user-item interaction. It consists of a user id, movie id and a resulting rating between 1 and 5, as well as a Unix timestamp of when the user-rating was logged. These represent the sparse features of the dataset. The other parts hold dense features about the user and the movies. The demographic data entails information about the user age, their gender, their occupation and the zip code. The movie data is made up of the movie name, its release date, a link to the IMDb[[10]](#footnote-11) page of the movie, and columns for each genre. The genres are encoded in a binary fashion, meaning that genres belonging to the movie are assigned the value 1, otherwise it’s set to 0.

### Data Preparation

All files prior to the pipeline ingestion are stored in the *data* directory. The raw dataset download files are stored in the *download* subdirectory. The files in this folder cannot be ingested by a MLOps pipeline yet and need to undergo data preparation steps. Usable data is stored in *recommender-datasets*. This folder serves as a repository for ingestible data. From there the datasets inside *recommender-datasets* are copied over to separate dedicated pipeline folders. There, the specific dataset is stored to be ingested by the respective MLOps pipeline (Figure 13).

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Figure 13: Structure of data directory

Having established the structure of the *data* directory, now the data preparation process is described, which uses the script movielens\_csv\_generator.ipynb.

First, the MovieLens 100k dataset is downloaded from the GroupLens file repository (GroupLens, n.d.). The dataset comes in a zip-file, which needs to be extracted. After the contents have been unpacked, the zip file is deleted. This process is done with the following inline commands in the Jupyter notebook:

# Fetch data (only run for the first time)

!wget https://files.grouplens.org/datasets/movielens/{ds\_name}.zip

!unzip {ds\_name}.zip

!rm {ds\_name}.zip

The extracted files are saved in the *downloads* folder. The dataset is divided into separate files. In order to create a complete dataset, the files need to be concatenated. For this research, the user demographics will be joined together with the sparse data of the user-item interaction. For this, *pandas* is used, a python library for data processing and data analysis (pandas, n.d.). The individual parts of the dataset are loaded into pandas and merged into one dataframe. Dataframes are objects that hold data in form of a table. Dataframes support a wide array of operations to transform and manipulate the data, which is the reason it is used for this data preparation task.

# Augment data\_df with user\_df

df = data\_df.merge(user\_df, on='user\_id')

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Figure 14: Sparse rating features and dense user features merged

The dataset used by the Recommender System will use 5 features, with the movie rating as the label:

* *user\_id* (sparse)
* *movie\_id* (sparse)
* *raw\_user\_age* (dense)
* *user\_gender* (dense)
* *user\_occupation* (dense)
* *user\_rating* (sparse, label)

Since SotA recommender systems can leverage both sparse and dense features to train a model, it was a priority to create a dataset that incorporates a mixture of both types of features. In order to simplify the data ingestion process for the RS, all features should be categorial and ordinal values. All features are already categorical besides *raw\_user\_age*, which is continuous. In order to make *raw\_user\_age* categorical, each user age is assigned to a specific age group, seen in Table 3. These age cohorts are derived from other MovieLens datasets (GroupLens, n.d.).

Table 3: Age cohorts

|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
| **Age group** | 1 | 18 | 25 | 35 | 45 | 50 | 56 |
| **Age range** | <18 | 18-24 | 25-34 | 35-44 | 45-49 | 50-55 | 55< |

In the next step all features are integer encoded. This is done to easily generate a category vocabulary for the RS, which is explained in the following chapter. In Figure 15 the final dataset can be seen.

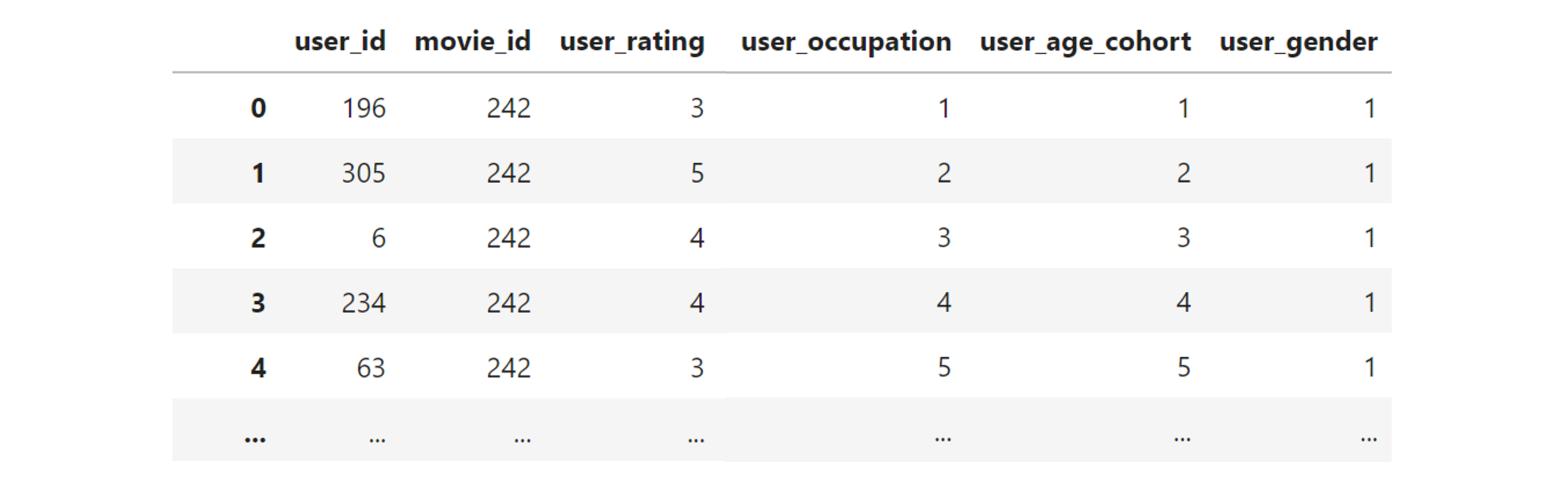


Figure 15: Pipeline dataset

After data preparation is done, the dataframe is saved in CSV format inside the *recommender-systems* folder. Throughout this project different data preparation approaches and data formats have been explored.[[11]](#footnote-12) Besides CSV, TensorFlow’s TFRecord format has been considered as well. Compared to CSV, it is more scalable and supports parallel-training in conjunction with TensorFlow. TFRecord is however only natively supported within the TensorFlow ecosystem. Ultimately, CSV has been chosen as the data format, because of its general compatibility with different tools and its ease of use with libraries like pandas. Since the relatively small MovieLens 100k dataset is used for this work, performance isn’t a detrimental factor in the selection of the file format.

## Recommender System

### Tools

### Implementation

**Data**.

**Recommender System**. The product of this component is *recommender\_system.py*.

### (Challenges)

## Concept Drift Awareness

## MLOps Pipeline

# Evaluation

# Conclusion

## Design Science Research

## Recommender Systems

### Overview

# Conclusion and Outlook

Mit der hier vorgelegten Dokumentvorlage steht ein Werkzeug zur Verfügung, das es möglich macht, auf relativ einfache Weise die Textgestaltung für Abschlussarbeiten und andere wissenschaftliche Arbeiten zu automatisieren. Dies spart Arbeitszeit, Beratungsaufwand und erhöht zugleich die Qualität des Ergebnisses hinsichtlich formaler Kriterien.

Gleichzeitig unterstützt die Dokumentvorlage die Lehre in Fächern *wie Arbeits-, Lern- und Präsentationstechniken* und *Vorbereitung auf die Bachelor- bzw. Masterarbeit*,

Die Dokumentvorlage wurde im Sommer 2000 in der vorliegenden grundlegend überarbeiteten Version zum ersten Mal hochschulweit benutzt, nachdem bereits die Vorgängerversion im Jahr 1999 für drei Abschlussarbeiten erfolgreich eingesetzt wurde.

Erweiterungen der Dokumentvorlage und der darin enthaltenen Anleitung zur Erstellung von Abschlussarbeiten mit dem Textsystem Microsoft Word sind auch für die Zukunft geplant. Das Feedback aus der Nutzung der Dokumentvorlage für Abschlussarbeiten an der HdM wird dabei fortgesetzt aufgegriffen und zur Verbesserung genutzt werden.

# Anhang A: Beispiele für die Gliederung von Abschlussarbeiten

Die nachfolgenden Gliederungen stellen lediglich Vorschläge dar, die stets am konkreten Fall überprüft und in der Regel angepasst werden müssen.

## A.1 Literaturarbeiten

1. Überblick (oder: Zusammenfassung, „Executive Summary“, alles Wichtige für den „Manager“ oder Schnellleser)
2. Fragestellung (oder: Ziele, Ausgangspunkt, Motivation)
3. Übersicht über den Stand der Wissenschaft und Technik (Beschreibung der Lösungsansätze, Beispiele etc. in einzelnen Abschnitten)
4. Bewertung der einzelnen untersuchten Ansätze, Beispiele etc., Identifikation von Defiziten
5. Synthese: Erstellung einer Gesamtschau, allgemeine Prinzipien, Beschreibung einer eigenen Sicht auf das Problem, evtl. auch eigene Vorschläge
6. Zusammenfassung (Erklärung des Nutzens), Ausblick

Anhang: eventuell recherchierte Texte, Produktbeschreibungen, etc.

## A.2 Systementwicklungen

1. Überblick (oder: Zusammenfassung, „Executive Summary“, alles Wichtige für den „Manager“ oder Schnellleser)
2. Problemstellung (oder: Ziele, Ausgangspunkt), Vorgesehener Benutzerkreis, Bedürfnisse der Benutzer
3. Stand der Technik (Wie wird das Problem bisher gelöst, wo sind die Defizite)
4. Gewählter Lösungsansatz (allgemeines Prinzip, welche Werkzeuge, z.B. Programmiersprachen werden verwendet)
5. Beschreibung der durchgeführten Arbeiten
6. Ergebnis (z.B. Screenshots mit Erläuterungen)
7. Zusammenfassung (Erklärung des Nutzens), Ausblick

Anhang: evtl. (ausgewählte) Programmbeispiele

Evtl. CD-ROM als Beilage

# Anhang B: Formatvorlagen

Nachfolgend sind die für die Benutzer der Dokumentvorlage wichtigsten Formatvorlagen aufgelistet. Vermerkt ist jeweils auch der Zweck der Formatvorlage und ob es sich um eine neue oder eine modifizierte Formatvorlage handelt.

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1. Throughout this paper ML and AI will be used interchangeably. [↑](#footnote-ref-2)
2. Rank refers to the dimensionality of a latent factor for Matrix and for Matrix , low-rank meaning that . represents the amount of linearly independent factors in a latent factor (Aggarwal (2016). [↑](#footnote-ref-3)
3. NNs and ANNs will be used interchangeably in this paper. [↑](#footnote-ref-4)
4. The Softmax function maps the neural output on range [0;1]. Highly negative inputs approach value 0, while highly positive inputs approach value 1. [↑](#footnote-ref-5)
5. In dropout regularization random neurons are skipped during the training process to combat overfit of the model (Hinton, Srivastava, Krizhevsky, Sutskever, and Salakhutdinov (2012). [↑](#footnote-ref-6)
6. Explicit feature interactions are usually retrieved by dedicated components in the model, such as the cross network in DCN. The order of feature interaction can be set (bounded-degree) and inferred by said component, which makes the feature extraction more controllable and predictable. On the contrary, implicit feature interactions can have an arbitrary order, as it is not explicitly defined in the model to which degree the interaction is bound to (i.e. DNN). Implicit feature extraction is more flexible at picking up any patterns in the data, at the cost of certainty of the degree of the feature interaction (Lian et al. (2018); Huang, She, Wang, and Zhang (2020); Yan and Li (2020). [↑](#footnote-ref-7)
7. A feature store is a data management layer that tracks and maintains features in ML data. Feature stores simplify ML pipelines and the data engineering process, by providing a centralized space where all raw and transformed data features are stored in. Previous data transformation steps are saved in the feature store, where they can be reused by other data scientists (Kakantousis et al. (2019). [↑](#footnote-ref-8)
8. Distribution and concept are used synonymously in the context of concept drift. [↑](#footnote-ref-9)
9. https://github.com/MyPetOctocat/bachelor\_2022 [↑](#footnote-ref-10)
10. IMDb (Internet Movie Database) is a database of entertainment media. It provides information about movies, TV shows and videogames. (IMDb, 2022) [↑](#footnote-ref-11)
11. The different data preparation approaches can be reviewed in the repository of this research under *artifact/data\_fetch* [↑](#footnote-ref-12)