

25 Key Equations in Machine Learning

Insight Hatch Machine Leanring Reviews - Vol 1

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Prefice and Introduction

This book is a journey through the 25 most important mathematical equations that underpin modern data science and machine learning. The idea is not only to learn about each of these equations but also to deeply understand their applications, background, and practical examples. We will explore each concept visually and instructively, shedding light on how these fundamental equations contribute to building intelligent systems.

Mathematics is the language of the universe, and it is the foundation of the incredible advances we see today in artificial intelligence and machine learning. The purpose of this book is to develop a learning guide that is both informative and visually compelling, bringing out the beauty and utility of these powerful mathematical tools. From optimization algorithms like Gradient Descent to classification techniques like Naive Bayes, and from measures of information like Entropy to clustering algorithms like K-Means, this book aims to provide an accessible yet thorough explanation of how these concepts work and why they are essential.

Each chapter focuses on one equation, starting with an introduction, followed by a detailed description, and ending with examples of how it is used in practice. Whether you're an aspiring data scientist, a seasoned engineer, or just curious about the mathematics that drives intelligent technology, this book will provide you with a solid understanding of these essential tools and how they interact to solve complex problems. Visual aids, illustrative examples, and in-depth explanations will help demystify each topic, making learning both engaging and enjoyable.

Each section will be followed by questions and homework problems. These problems are selected based on their popularity and effectiveness in enhancing understanding. The goal is to provide exercises that solidify the reader's comprehension of each topic. An answer key is provided at the end of the book to facilitate learning and self-assessment.

Many of the examples and problems use Python and R to provide practical insights. Code blocks in Python and R (and in some instances Mathematica) are used throughout to illustrate the concepts discussed. All the example codes, as well as the entire book, are posted on the author's GitHub repository for easy access and reproducibility.

Whenever possible, all examples that originate from a particular source are acknowledged by citation and recognition of the original author. If any citation or attribution is missed, the author kindly requests feedback so that proper corrections can be made.

Some notes on the typesetting framework and tools

With the current advent of ML based LLMs and the availability of typesetting LaTeX, we combine tools like Mathpix (the LaTeXequation generating snipping tool) and R Markdown driven by Pandoc and knitr to essentially "GET IT DONE". Obviously your mileage may vary; we find this tool set perfect for typesetting combined with computational analysis. Why? Well you have nearly full LaTeXcapability with access to R & Python (via reticulate) programming code blocks and also the ability to bridge to MATHEMATICA® if needed. On a powerful laptop, this tool suite (under RStudio) leaves the door open to a wider world of analysis and computation. While RStudio is not perfect, it does allow for an IDE approach which leverages typesetting and computation framework (through R, Python and MATHEMATICA®) that makes it a worthy Swiss Army Knife. The alternative is also VS Code, which is great, however the mixed mode R Markdown & LaTeXcomputational environment is pretty flexible.

The complimentary nod to ML LLM tools

Most of this compilation and summary work would not be possible without the advent of LLMs and new found ability to build concise summaries of difficult mathematical concepts. The time savings is basically a gift which save more grey hairs and time flipping pages of numerous cross references. Actually we want to spend time "learning" and not "flipping and skimming" to find the properly defined statement which is of utility in our endeavor. So for that - we lean on LLMs to be more productive, and help clearly convey underlying concepts. This is not to be lazy, each section must be scrutinized for accuracy and consistency. The speed gains allow for a very large group of examples and highly detailed presentations.

Notes to usage

We give full credit for any original works that are used herein to reach the stated objective. At this point this is not a financial pursuit we do not provide any license related respect to any previous publisher. Go figure. Why would we take such a stand - this is educational open source material. We strongly recommend you take the time to do some background research on any referenced authors.

We truly hope you enjoy the tour, as much as we have enjoyed our journey to write it... to-ohru iwanami, 2024, somewhere in asia

The 25 (uhh 24)

How did we arrive at the 25 (uhh 24) most important equations? Well, it's partly a matter of educated opinion, partly a rough statistical average of what people talk about most when they're exhilarated by the magic of machine learning, and partly—let's be real, you should just listen to me (fully joking here). These equations are more than just mathematical tools; they represent milestones that have shaped the evolution of artificial intelligence, each one contributing a brick or building block in the formidable wall of progress we've built over the decades.

Take, for instance, Gradient Descent — a cornerstone of optimization, whose principles date back to the early 19th century and the work of Adrien-Marie Legendre and Carl Friedrich Gauss in minimizing residuals. The modern incarnation of Gradient Descent, essential for training deep neural networks, only emerged in the mid-20th century, revitalized by researchers trying to teach machines to "learn." Fast forward to the 1960s, when Frank Rosenblatt was developing the perceptron, it became evident that iterative optimization methods like Gradient Descent could unlock the potential of early neural networks.

Then we have Bayes' theorem, after Reverend Thomas Bayes, who developed his idea of conditional probability back in the 18th century. Although it lay relatively dormant for years, it became a key breakthrough in the 1950s when Alan Turing and others applied Bayesian methods to codebreaking during World War II. Today, its cousin—Naive Bayes — is still a powerful tool for classification, especially for natural language processing, allowing us to create chatbots and email spam filters. This dormant theorem found new life, propelling us toward a world of probabilistic understanding in machines.

Singular Value Decomposition (SVD), another member of our elite list, found its fame in the 1990s when it was used for information retrieval in the famous Latent Semantic Analysis (LSA) algorithm—paving the way for how we handle and understand textual data. The underlying mathematics, discovered by Eugenio Beltrami and others in the 19th century, helps us not only reduce dimensionality but also reveal the hidden relationships between concepts in data.

Fast forward to today to the ReLU (Rectified Linear Unit) function. While it seems so simple,

this piece of mathematical elegance emerged as a game-changer for deep learning in the 2010s, thanks to the work of Geoffrey Hinton and his colleagues. Before ReLU, neural networks struggled with the vanishing gradient problem, limiting their ability to learn effectively. By simply transforming negative inputs to zero and retaining positive ones, ReLU helped neural networks go deeper and deeper (pun-ch!), giving birth to the deep learning revolution we know today.

Each of these equations is a testament to the evolution of knowledge, a story of breakthroughs spanning centuries. Together, they form a bridge from the deterministic calculus of Newton and Leibniz to the probabilistic dreams of Bayes and the practical algorithms of today's AI pioneers. Think of them as the greatest hits of mathematics for data scientists—carefully curated, occasionally debated, and ultimately distilled into the essence of what drives machine learning today. This collection captures a human legacy of curiosity and ingenuity, guiding you on your intellectual journey through the ever-growing forest of machine learning—one equation, one insight at a time.

As we reflect on these 1 through 24 equations, and on to 25, one cannot ignore the human spirit that threads through each mathematical symbol and formula. Behind every breakthrough is a mind driven by curiosity, a determination to see beyond the obvious, to turn abstraction into discovery. Each equation, in its own way, captures a moment where human thought transcended barriers—whether it was a problem of optimization, uncertainty, or understanding the nature of learning itself. Yet, even as these 24 pillars of insight stand tall, there remains one more — a keystone that binds them all, born not out of calculation alone but from the essence of what it means to explore, to question, and to create. This final entry goes beyond numbers and symbols, embracing the very source of every theorem, every insight, and every spark of ingenuity. So as you journey onward, remember—there's always one more equation to unveil, one that's been with us all along.

Equation 1

Gradient Descent

$$\theta_{j+1} = \theta_j - \alpha \nabla J(\theta_j)$$



Finding a local minimum with Gradient Descent

Key ML Equation 1: Gradient Descent

$$\boldsymbol{\theta}_{i+1} = \boldsymbol{\theta}_i - \alpha \nabla J(\boldsymbol{\theta}_i) \tag{1.0.1}$$

- θ_j The current value of the **parameter vector** at iteration j, which represents the current estimate of the model parameters.
- θ_{j+1} The updated value of the **parameter vector** for the next iteration, which results from applying the gradient step to the current parameters.
- α The **learning rate**, which is a positive scalar determining the size of the step to take in the direction of the negative gradient.
- J The **cost function**, which is the function being minimized by adjusting the parameter vector θ . It measures the difference between predicted values and actual values.

Introduction: Gradient Descent is an optimization algorithm used to minimize the cost function by iteratively moving in the direction of the steepest descent as defined by the negative of the gradient.

Description: In machine learning, gradient descent is used to update the parameters of the model, θ , to reduce the difference between the predicted and actual outcomes.

Importance in ML: Gradient Descent is foundational for training machine learning models, particularly in neural networks and linear regression. It helps in finding optimal parameters by iteratively reducing the error, making it crucial for model accuracy.

1.1 What is behind the equation

1.2 Structure of the Parameter Vector θ

The parameter vector, typically denoted as θ , contains all the adjustable parameters or weights of the model that you want to optimize in order to minimize the cost function $J(\theta)$. Depending on the type of model, the structure of θ can vary:

• Linear Regression:

$$\theta = \begin{bmatrix} \theta_0 \\ \theta_1 \\ \vdots \\ \theta_n \end{bmatrix} \tag{1.2.1}$$

In linear regression, θ is often a column vector of weights where each θ_i corresponds to the weight associated with feature x_i . θ_0 is often referred to as the bias term or intercept.

• Logistic Regression / Neural Networks: In these models, θ can have more dimensions. For a neural network, the parameter vector could be a collection of weight matrices for different layers, such as:

$$\theta = \{W_1, W_2, \dots, W_L, b_1, b_2, \dots, b_L\} \tag{1.2.2}$$

where W_i and b_i are weights and biases associated with layer i of the network.

• Deep Learning Models: The parameter vector is much more complex, often containing multiple matrices and vectors representing weights and biases for each layer in a deep neural network.

In general, θ is a vector that can be expressed as:

$$\boldsymbol{\theta} = (\theta_1, \theta_2, \dots, \theta_n)^T \tag{1.2.3}$$

where n is the number of features or neurons, depending on the model type.

1.3 Typical Value of the Learning Rate α

The learning rate α controls the step size when updating the parameters during gradient descent. Choosing a proper value for α is crucial for the convergence of the algorithm. Here are some general guidelines:

• **Typical Range**: A typical value for the learning rate lies between 0.001 and 0.1. Values outside this range can either lead to a slow convergence or an unstable training process.

• Considerations:

- **Too Small**: If α is too small, gradient descent will take very small steps towards the optimum, resulting in a very slow convergence process.
- **Too Large**: If α is too large, gradient descent might overshoot the minimum or even diverge, causing the cost function to oscillate or increase.
- Adaptive Learning Rates: Some advanced algorithms like Adam use adaptive learning rates which adjust automatically as training progresses.

In practice, tuning the learning rate often involves trial and error or the use of techniques like **learning rate schedules** or **grid search** to find the best value for a specific problem.

1.4 Math Behind the Gradient $\nabla J(\theta_i)$

The term $\nabla J(\theta_j)$ is the **gradient** of the cost function $J(\theta)$ evaluated at θ_j . Let's break it down:

• Gradient Definition: The gradient of a function is a vector of partial derivatives with respect to each parameter in θ . In the case of $J(\theta)$, the gradient $\nabla J(\theta_j)$ tells us how much the cost function changes when we make an infinitesimally small change to each component of θ . Mathematically, for a parameter vector $\theta_j = (\theta_1, \theta_2, \dots, \theta_n)^T$:

$$\nabla J(\theta_j) = \begin{bmatrix} \left. \frac{\partial J}{\partial \theta_1} \right|_{\theta_j} \\ \left. \frac{\partial J}{\partial \theta_2} \right|_{\theta_j} \\ \vdots \\ \left. \frac{\partial J}{\partial \theta_n} \right|_{\theta_j} \end{bmatrix}$$

Each partial derivative $\frac{\partial J}{\partial \theta_i}$ represents the rate of change of the cost function with respect to parameter θ_i .

• Intuition: The gradient $\nabla J(\theta_i)$ points in the direction of the steepest ascent of the cost function J. In gradient descent, we want to **minimize** $J(\theta)$, so we move in the opposite direction, which is why the update rule is:

$$\theta_{j+1} = \theta_j - \alpha \nabla J(\theta_j)$$

- $\nabla J(\theta_j)$ represents the slope or direction in which $J(\theta)$ increases most quickly.
- By subtracting $\alpha \nabla J(\theta_j)$, we effectively move in the direction of steepest **descent**, hence reducing $J(\theta)$.
- Computing the Gradient: In practice, the gradient $\nabla J(\theta_i)$ is computed using differentiation. For different cost functions, the gradient takes different forms:
 - For linear regression with a mean squared error (MSE) cost function, the gradient is relatively simple and involves the residuals (errors) between predictions and actual values.
 - For **neural networks**, computing the gradient involves **backpropagation**, which is a process of applying the chain rule of calculus to calculate the gradients efficiently for each layer.

In short, $\nabla J(\theta_j)$ gives us the necessary information to adjust θ in a way that reduces the error. The learning rate α then determines how big the step should be in this direction.

1.5 Gradient Descent in R

Gradient Descent is a foundational optimization algorithm used to iteratively minimize cost functions. Here, we apply the gradient descent method to find the local minimum of a specific polynomial function. The function used is a quartic polynomial $P(x) = x^4 - 6x^3 + 11x^2 - 6x$. Our goal is to observe how the gradient descent algorithm updates our parameter over successive iterations to converge towards a minimum point of the polynomial.

The following R code demonstrates how to implement gradient descent for this polynomial, including plotting the descent path and function value across iterations. This practical example aims to give you a clear understanding of how gradient descent operates on real functions, using R for illustration. Note that gd_result is the dataframe result fed to ggplot.

```
# Set seed for reproducibility
set.seed(42)
\# Define the new polynomial function and its derivative
polynomial_function <- function(x) {</pre>
  return(x^4 - 8 * x^3 + 18 * x^2 - 11 * x + 2)
# Derivative of the new polynomial function
polynomial_derivative <- function(x) {</pre>
  return(4 * x^3 - 24 * x^2 + 36 * x - 11)
}
# Gradient Descent Algorithm
gradient_descent <- function(learning_rate = 0.01, iterations = 1000, start = 3) {</pre>
  x <- start # Starting point
  # Store x values and function values for plotting
  x_values <- numeric(iterations)</pre>
  function values <- numeric(iterations)</pre>
  for (i in 1:iterations) {
    grad <- polynomial_derivative(x)</pre>
    x <- x - learning_rate * grad
    x_values[i] <- x</pre>
    function_values[i] <- polynomial_function(x)</pre>
  return(data.frame(Iteration = 1:iterations, x_values, function_values))
}
# Run the Gradient Descent with specific parameters
learning_rate <- 0.01</pre>
iterations <- 100
start_point <- 3
gd_result <- gradient_descent(learning_rate, iterations, start_point)</pre>
```

The following R code demonstrates how to implement gradient descent for this polynomial, including plotting the descent path and function value across iterations. This practical

example aims to give you a clear understanding of how gradient descent operates on real functions, using R for illustration.

```
library(ggplot2)
ggplot(gd_result, aes(x = Iteration, y = function_values)) +
    geom_line(color = "blue", size = 1.2) +
    ggtitle("Gradient Descent on Polynomial Function") +
    xlab("Iteration") +
    ylab("Objective Function Value: f(x)") +
    theme_minimal() +
    theme(
        plot.title = element_text(size = 11)  # Adjust the size value as needed
)
```

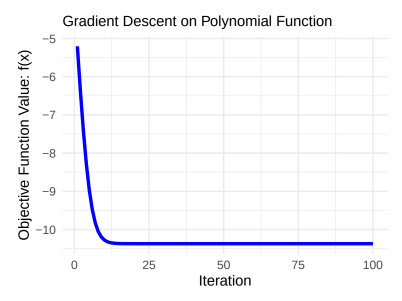


Figure 1.1: Gradient Descent on Polynomial Function - Iteration vs Function Value

```
# Additional Plot: Trace the Path of Gradient Descent on the Polynomial
x_range <- seq(min(gd_result$x_values) - 1, max(gd_result$x_values) + 1, length.out = 500)
polynomial_values <- polynomial_function(x_range)

ggplot() +
    geom_line(aes(x = x_range, y = polynomial_values), color = "black", size = 1) +
    geom_point(data = gd_result, aes(x = x_values, y = function_values), color = "red", size = 1.5) +
    ggtitle("Gradient Descent Path on Polynomial Function") +
    xlab("x") +
    ylab("f(x)") +
    theme_minimal() +
    theme(
        plot.title = element_text(size = 11) # Adjust the size value as needed
    )</pre>
```

Gradient Descent Path on Polynomial Function

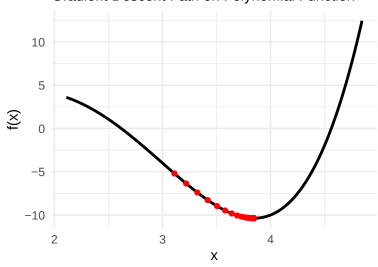


Figure 1.2: Gradient Descent on Polynomial Function - Trace the Path on the Polynomial

1.6 Linking Gradient Descent to Our Polynomial Example

The equation we used to explain gradient descent is:

$$\theta_{i+1} = \theta_i - \alpha \nabla J(\theta_i)$$

1.6.1 Step-by-Step Breakdown

The fundamental idea behind gradient descent is to iteratively adjust the model's parameters, θ , in a direction that reduces the cost function, $J(\theta)$. The parameter update is governed by the **gradient** of the cost function (or loss function), $\nabla J(\theta)$.

In the above equation:

- θ_j represents the parameter(s) at step j, and θ_{j+1} is the updated parameter at step j+1.
- α is the **learning rate**, which controls how large each step is in the descent.
- $\nabla J(\theta_j)$ is the **gradient** of the cost function with respect to the parameters, evaluated at θ_j . It gives the direction of the steepest ascent, and since we want to minimize the function, we move in the opposite direction, hence the negative sign.

1.6.2 Gradient Descent Applied to the Polynomial

In our R code example, we have a **polynomial function** defined as:

$$P(x) = x^4 - 6x^3 + 11x^2 - 6x$$

Our goal is to minimize this polynomial function, which represents our **cost function**, J(x). Here, x plays the role of our parameter, θ , that we want to adjust iteratively using gradient descent.

The **derivative** of the polynomial is:

$$P'(x) = 4x^3 - 18x^2 + 22x - 6$$

In the gradient descent algorithm, this derivative represents the **gradient** of the cost function with respect to our parameter, x. This means that $\nabla J(x) = P'(x)$. The iterative update step becomes:

$$x_{j+1} = x_j - \alpha P'(x_j)$$

Where: - x_j is the current value of the parameter (similar to θ_j). - α is the learning rate that we set in the R code. - $P'(x_j)$ is the gradient of the polynomial at the current point.

1.6.3 Bringing It Full Circle

In our R code, we used **gradient descent** to minimize the polynomial function by starting at an initial point (x = 3) and repeatedly updating x using the gradient descent rule:

This corresponds exactly to the equation:

$$x_{j+1} = x_j - \alpha P'(x_j)$$

As each iteration proceeds, x gets closer and closer to a point where the gradient is zero (i.e., a local minimum or a stationary point). In this particular case, the polynomial P(x) has one real minimum, and the gradient descent algorithm helps us converge towards it.

1.6.4 Visual Connection

The **first plot** in our example (function value vs. iteration) shows how the value of the polynomial decreases over time as gradient descent proceeds. The **second plot** (gradient descent path on the polynomial curve) visually shows how x moves along the polynomial curve, getting closer and closer to the minimum.

This linkage between the theoretical equation of gradient descent and the practical implementation of minimizing the polynomial demonstrates how gradient descent serves as a universal tool to solve optimization problems, whether in machine learning or even in simple polynomial functions like the one used in our example.

Gradient descent works by always taking steps in the direction that most rapidly reduces the cost function—allowing us to find optimal parameters, minimize errors, or reach local minima. It is an essential part of training many machine learning models, and the concept remains consistent regardless of the specific problem context.

1.7 Additional ideas to explore

Stochastic Gradient Descent (SGD), Mini-Batch Gradient Descent, or concepts like Momentum and Learning Rate Scheduling.

We could also explore more code examples, such as:

Adding stopping criteria to our gradient descent R code, like a tolerance for change in cost function.

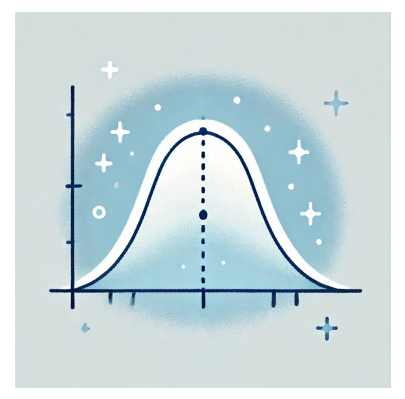
Visualizing the convergence of the gradient descent with multiple starting points.

Implementing different cost functions and comparing their optimization trajectories. Let me know which direction you'd like to explore, and I'll get started!

Equation 2

Normal Distribution

$$f(x|\mu,\sigma^2) = \frac{1}{\sigma\sqrt{2\pi}} \exp\left(-\frac{(x-\mu)^2}{2\sigma^2}\right)$$



Seeing the average and variance with the Normal Distribution

Key ML Equation 2: Normal Distribution

$$f(x|\mu,\sigma^2) = \frac{1}{\sigma\sqrt{2\pi}} \exp\left(-\frac{(x-\mu)^2}{2\sigma^2}\right)$$
 (2.0.1)

- The **random variable** for which the probability density function is being calculated. It represents the value within the distribution.
- μ The **mean** of the distribution, which represents the center or "average" value around which the data clusters.
- σ The **standard deviation** of the distribution, indicating how spread out the values are around the mean. A larger σ means more spread, while a smaller σ means values are more tightly clustered.
- σ^2 The **variance** of the distribution, which is the square of the standard deviation. It provides a measure of the dispersion of the distribution.
- $f(x|\mu, \sigma^2)$ The **probability density function (PDF)** for the normal distribution, which provides the likelihood of x occurring given the parameters μ and σ^2 .
- exp The **exponential function**, which ensures that the PDF value falls off symmetrically from the mean μ . It plays a key role in modeling the bell-shaped curve characteristic of the normal distribution.

Introduction: The normal distribution, also called the Gaussian distribution, is a probability distribution that is symmetric about the mean.

Description: It represents how data tends to cluster around a central point. The parameters μ and σ^2 represent the mean and variance, respectively.

Importance in ML: The normal distribution is used in many ML algorithms, especially in probabilistic models and hypothesis testing. Assumptions of normality often simplify the mathematics of learning models and are vital in Bayesian networks.

2.1 What is Behind the Equation

The normal distribution equation represents a probability density function (PDF) that models how data points are distributed around a central value (the mean, μ). This equation is essential in statistics because it describes a common pattern found in natural phenomena, from heights of people to errors in measurements. The bell-shaped curve is a striking visual, representing how values tend to cluster around the mean, with fewer observations occurring as we move further from this center. The beauty of the normal distribution lies in its symmetry and the way it characterizes many real-world datasets, making it a cornerstone in both statistics and machine learning.

2.2 The Players

Normal distribution

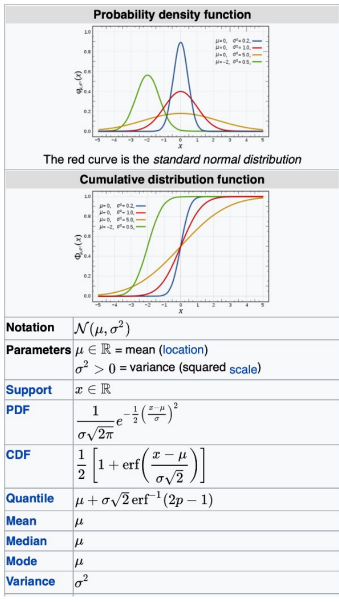


Figure 2.1: Summary of Normal Distribution with PDF and CDF



Figure 2.2: Carl Friedrich Gauss



Figure 2.3: Pierre-Simon Laplace

- 1. Normal Distribution: [Normal Distribution on Wikipedia]
- 2. Carl Friedrich Gauss: [Carl Friedrich Gauss on Wikipedia]
- 3. Pierre-Simon Laplace: [Pierre-Simon Laplace on Wikipedia]

2.3 Historical Context

The normal distribution, often referred to as the Gaussian distribution, has a rich history rooted in the development of probability theory and statistics. The distribution is named after the German mathematician Carl Friedrich Gauss, who used it extensively in his work on astronomy and measurement errors in the early 19th century. Gauss formalized the idea that errors in measurements tend to follow a symmetric pattern around the true value, leading to the bell-shaped curve we now associate with the normal distribution.

However, the concept of the normal distribution predates Gauss and can be traced back to the work of Abraham de Moivre, an 18th-century French mathematician. De Moivre first derived the normal distribution as an approximation to the binomial distribution when the number of trials becomes very large. His work laid the foundation for what would later be formalized and popularized by Gauss.

The normal distribution became particularly significant due to the Central Limit Theorem, which states that the sum of many independent random variables, regardless of their original distribution, tends to follow a normal distribution. This theorem, proven by mathematicians such as Pierre-Simon Laplace, helped establish the normal distribution as a fundamental tool in statistics and the natural sciences. Today, it is widely used not only because of its mathematical properties but also because it naturally arises in numerous real-world situations, making it one of the most important and recognizable distributions in probability and statistics.

2.4 Relative Standard Deviation (RSD)

In the context of the normal distribution, the spread of data points is characterized by the standard deviation (σ). A useful concept derived from this is the **Relative Standard Deviation (RSD)**, which is expressed as a percentage of the mean:

$$RSD = \left(\frac{\sigma}{\mu}\right) \times 100$$

Typical RSD Values: The RSD provides insight into how variable the data is relative to its mean. A low RSD (e.g., below 10%) indicates that the data points are closely clustered around the mean, whereas a higher RSD suggests more dispersion. In many real-world datasets, RSDs ranging between 5% and 20% are common, depending on the type of measurement and its inherent variability. RSD helps give a standardized view of spread that is independent of the scale of the data, which is particularly useful when comparing the variability between datasets.

2.5 Understanding the Notation $f(x|\mu, \sigma^2)$

The notation $f(x|\mu, \sigma^2)$ represents the **probability density function** of a normal distribution given the parameters μ (mean) and σ^2 (variance). This notation indicates a **conditional** relationship, where the probability density function f(x) is dependent on the parameters μ and σ^2 .

In other words, μ and σ^2 define the specific characteristics of the distribution (where it is centered and how spread out it is), while x represents the variable for which the probability is being calculated. This notation highlights that the distribution is characterized by these parameters, and the output f(x) tells us how likely it is to observe a particular value of x under that distribution.

2.6 The Nature of the Exponential Function in Normal Distribution

A key component of the normal distribution equation is the **exponential function**:

$$\exp\left(-\frac{(x-\mu)^2}{2\sigma^2}\right)$$

This part of the function gives the normal distribution its famous **bell shape**. The negative squared term in the exponent ensures that values closer to the mean (μ) have higher probabilities, while values further away have exponentially decreasing probabilities. This shape is what makes the normal distribution symmetric, with a single peak at the mean.

The function $\exp(-x^2)$ falls off rapidly as |x| increases, which results in a smooth, continuous decline from the peak at the mean. This property is crucial because it reflects how, in many natural phenomena, values tend to cluster around an average, with extreme values being much less common. This elegant behavior is what makes the normal distribution so special and why it is so frequently used in statistical modeling.

2.7 Non-existence of an Analytical Anti-Derivative

Interestingly, the normal distribution function does not have an anti-derivative that can be expressed in closed form. This means that the area under the curve (which represents the cumulative probability) cannot be solved using traditional analytic integration techniques. Instead, it is computed numerically or looked up using statistical tables.

The integral of the normal distribution is given by:

$$\int_{-\infty}^{\infty} f(x|\mu, \sigma^2) \, dx = 1$$

However, there is no elementary function that represents this integral. This is why the **error** function (erf) is introduced in mathematics to help approximate these values:

$$\int e^{-x^2} dx = \frac{\sqrt{\pi}}{2} \operatorname{erf}(x) + C$$

The lack of an analytical solution means that in practice, probabilities for a normal distribution are often calculated using **numerical methods** or **precomputed tables**. This characteristic of the normal distribution makes it an interesting function from a mathematical standpoint, as it combines simplicity of form with complexity in integration.

2.8 Central Limit Theorem

The Central Limit Theorem (CLT) is a fundamental concept in probability theory that explains why the normal distribution is so prevalent in nature and in machine learning. The CLT states that when independent random variables are added together, their properly normalized sum tends to form a normal distribution, regardless of the original distribution of the variables. This remarkable property applies as long as the number of variables is sufficiently large and they have finite variances.

Mathematically, if we have a set of random variables, each with mean and variance, the sum or average of these variables will tend towards a normal distribution as becomes large. The formal expression is:

where denotes convergence in distribution, and represents the standard normal distribution.

The CLT helps explain why the normal distribution is observed so often in practice. Many complex processes can be thought of as the sum of many small, independent effects. Whether it's measurement errors, human heights, or even financial market fluctuations, these effects combine to form a distribution that is approximately normal.

2.8.1 Importance in Machine Learning

The Central Limit Theorem has significant implications in machine learning. It provides the foundation for many statistical techniques and justifies the assumption of normality in models. For example:

Modeling Errors: In regression analysis, the residuals (errors) are often assumed to be normally distributed. This assumption allows us to derive confidence intervals and perform hypothesis

testing.

Feature Engineering: When aggregating data, such as calculating the mean of multiple features or observations, the resulting values tend to be normally distributed, making it easier to apply techniques that assume normality.

Sampling Distributions: The CLT allows us to make inferences about population parameters from sample data. Many machine learning algorithms rely on sampling and estimation, and the CLT ensures that the distribution of the sample mean approximates normality, which simplifies analysis and interpretation.

The CLT thus serves as a bridge between randomness and order, allowing us to apply the powerful tools of the normal distribution even in cases where the underlying data might not be normally distributed. It is this ability to generalize and predict outcomes that makes the normal distribution so central to both statistics and machine learning.

2.9 Applications in Machine Learning

The normal distribution plays a vital role in numerous machine learning algorithms and concepts. Its presence is seen in everything from model assumptions to data transformations. Below are some of the key areas where the normal distribution is commonly applied:

1. Gaussian Naive Bayes

Gaussian Naive Bayes is a classification algorithm based on Bayes' theorem. It assumes that the features follow a normal distribution, which allows the model to calculate probabilities efficiently. The assumption of normality simplifies the calculations, enabling rapid classification even with high-dimensional data. This assumption works well for many real-world datasets, making Gaussian Naive Bayes a popular choice for problems like spam detection and document classification.

2. Linear Regression Error Terms

In linear regression, the error terms (residuals) are often assumed to be normally distributed. This assumption allows us to make statistical inferences about the parameters of the regression model, such as constructing confidence intervals and conducting hypothesis tests. If the residuals are approximately normal, we can apply powerful statistical tools to evaluate model fit and make predictions.

3. Weight Initialization in Neural Networks

When training neural networks, the weights are often initialized using a normal distribution. For example, in the Xavier initialization method, weights are drawn from a normal distribution with a mean of zero and a variance that depends on the number of input and output nodes.

This helps ensure that the neurons start with a diverse range of values, which prevents issues such as all neurons producing the same output. Proper initialization is crucial for efficient training, and using the normal distribution helps keep the gradients within a reasonable range during backpropagation.

4. Generative Models

The normal distribution is also used in generative models, such as Gaussian Mixture Models (GMMs), which assume that the data is generated from a mixture of several Gaussian distributions. GMMs are widely used in clustering problems, where they help to model the underlying distribution of the data and assign probabilities to different clusters.

5. Feature Scaling and Data Transformation

In many machine learning algorithms, it is beneficial for features to follow a normal distribution. Methods such as StandardScaler in scikit-learn standardize features by removing the mean and scaling to unit variance, resulting in a distribution with a mean of 0 and a standard deviation of 1. This process is especially important for algorithms that are sensitive to the scale of the input features, such as support vector machines (SVMs) and gradient descent optimization.

Importance of the Normal Distribution in Machine Learning

The normal distribution is not only a convenient assumption but also a useful tool in machine learning. Its prevalence in real-world phenomena and its mathematical properties make it indispensable for building, evaluating, and optimizing models. Whether it's in simplifying calculations through Gaussian Naive Bayes, making statistical inferences in linear regression, initializing neural networks effectively, or clustering data in GMMs, the normal distribution is at the core of numerous machine learning practices. By understanding and leveraging the normal distribution, practitioners can better model uncertainties and improve the robustness of their machine learning solutions.

2.10 Suggested Additional Content for the Chapter

- 1. **Historical Context**: Add a brief history of the normal distribution, perhaps mentioning Carl Friedrich Gauss, who contributed to its development, and why it is often called the Gaussian distribution.
- 2. **Visualization**: Add a visualization of the normal distribution with varying means and standard deviations to help illustrate how changes in μ and σ affect the shape of the curve.

Equation 3

Z-Score

$$z = \frac{x - \mu}{\sigma}$$



Take a portion of the probability with the Z-Score

Key ML Equation 3: Z-Score

$$z = \frac{x - \mu}{\sigma} \tag{3.0.1}$$

- The **random variable**, representing the value within the distribution that is being standardized.
- μ The **mean** of the distribution, which is the average value and the point around which data tends to cluster.
- σ The **standard deviation** of the distribution, indicating the spread or dispersion of values around the mean. A larger σ suggests more variability.
- The **Z-Score**, which represents how many standard deviations a particular x value is from the mean μ . It is a measure of relative position within the distribution.

Introduction: The Z-score represents the number of standard deviations a data point is from the mean.

Description: It is used to standardize data points within a dataset, making comparisons between different distributions possible.

Importance in ML: Z-scores are crucial in feature scaling and normalization, allowing different features to be compared and helping gradient-based algorithms converge faster by ensuring all features have a similar scale.

What is Behind the Equation

The Z-score, also known as the standard score, is a measure that describes the position of a value relative to the mean of a dataset, in units of the standard deviation. The Z-score calculation is particularly useful in the context of the normal distribution, allowing us to determine how far a particular value x lies from the mean μ when expressed in terms of the distribution's standard deviation σ . The Z-score is expressed by the formula:

$$z = \frac{x - \mu}{\sigma}$$

This metric is instrumental in transforming individual data points into a universal scale, where positive Z-scores indicate values above the mean, and negative Z-scores indicate values below the mean. The Z-score effectively normalizes different datasets, making comparisons straightforward.

3.1 Applications in Machine Learning

In machine learning, Z-scores are employed for several purposes, ranging from outlier detection to feature scaling.

- Outlier Detection: Z-scores can help identify outliers, as values with extremely high or low Z-scores typically indicate data points that lie far from the distribution's average behavior. These outliers might signify anomalies or valuable insights that need closer examination.
- Feature Scaling: Z-score normalization, also known as standardization, is a common feature scaling method. By transforming features using Z-scores, they are rescaled to have a mean of 0 and a standard deviation of 1. This type of normalization is especially valuable in algorithms like logistic regression, k-means clustering, and principal component analysis (PCA), where features with different ranges can negatively impact model performance.
- Standard Normal Table Applications: The Z-score is also used in combination with the standard normal distribution table to compute probabilities and p-values for hypothesis testing. In machine learning, this is often relevant in model evaluation and statistical testing.

3.2 Typical Range for Z-Scores

The typical range for Z-scores in a standard normal distribution is between -3 and 3. Values beyond this range are considered rare and are often associated with outliers.

- |Z| > 3: Typically considered outliers. Values beyond three standard deviations are quite unusual in a normal distribution.
- |Z| < 1: Most values are likely within this range and close to the mean.
- $1 \le |Z| < 2$: These values are still common, although they are further away from the mean.
- |Z| > 2: Values in this range start to become unusual, indicating either naturally rare observations or data quality issues.

The calculation of Z-score can be a valuable indicator of how far a data point deviates from what is expected, helping differentiate between normal variance and potential outliers.

3.3 Mathematical Notation and Concept

The equation for the Z-score is represented as follows:

$$z = \frac{x - \mu}{\sigma}$$

- x: Represents the actual observed value within the dataset.
- μ : The mean of the dataset, which represents the central tendency around which the dataset is distributed.
- σ : The standard deviation, which gives insight into the degree of variability within the data points. It tells us how spread out the values are around the mean.

The Z-score essentially tells us how many standard deviations away from the mean a given value x is, and whether it is to the left (negative z) or right (positive z) of the mean.

3.4 Mathematical Insights: Area Under the Normal Curve

One of the key properties of the Z-score is how it allows for the calculation of probabilities from the normal distribution. By converting a value to its Z-score, we can use the standard normal distribution (which has a mean of 0 and a standard deviation of 1) to determine the probability that a value is less than or greater than a given point.

For example, calculating the probability $P(Z \leq z)$ involves integrating the probability density

function of the normal distribution up to the given Z-score value:

$$P(Z \le z) = \int_{-\infty}^{z} \frac{1}{\sqrt{2\pi}} e^{-t^2/2} dt$$

Since this integral has no closed-form solution, it is typically evaluated using numerical methods, standard normal distribution tables, or software. This integral defines the cumulative distribution function (CDF) of the normal distribution, which is essential in assessing cumulative probabilities for standard normal variables.

In practice, machine learning models often use pre-calculated tables or libraries to compute these probabilities for performance metrics, hypothesis testing, or evaluating the significance of model parameters.

3.5 Example: Z-Score in R

The Z-score calculation can be easily implemented in R to standardize a dataset or to identify outliers. Below is a simple example to illustrate how we can compute Z-scores for a given dataset and visualize the data distribution using a histogram:

```
# Load required libraries
library(ggplot2)
# Define the parameters for the normal distribution
mean_value <- 0
std_dev <- 1
z score <- 1.25
# Define the cumulative distribution function (CDF) to calculate probability
p_value <- pnorm(z_score, mean = mean_value, sd = std_dev)</pre>
# Generate data for plotting the normal distribution curve
x_values \leftarrow seq(-4, 4, length.out = 1000)
y_values <- dnorm(x_values, mean = mean_value, sd = std_dev)</pre>
data <- data.frame(x = x_values, y = y_values)</pre>
# Plot the normal distribution curve with shaded area under the curve
plot <- ggplot(data, aes(x = x, y = y)) +
  geom_line(color = "black", size = 1) +
  geom_area(data = subset(data, x <= z_score), aes(x = x, y = y), fill = "steelblue", alpha = 0.5) +</pre>
  geom_vline(xintercept = z_score, color = "red", linetype = "dashed", size = 1) +
  ggtitle("Normal Distribution with Z-Score of 1.25") +
 xlab("Z") +
  ylab("Density") +
  theme_minimal()
# Print the plot
print(plot)
```

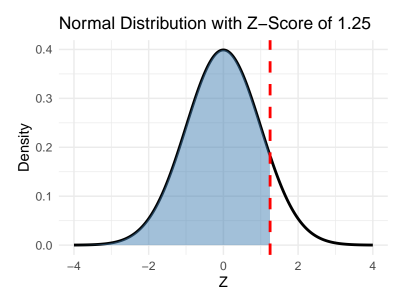


Figure 3.1: Default caption for all figures

```
# Print out the results
cat("Z-Score:", z_score, "\n")

## Z-Score: 1.25
cat("Probability (P-value) for Z <=", z_score, ":", round(p_value * 100, 2), "%\n")</pre>
```

Probability (P-value) for Z <= 1.25 : 89.44 %

In this code block: * We first generate a dataset of 100 normally distributed random numbers with a mean of 50 and a standard deviation of 10. * We calculate the Z-scores for each value by subtracting the mean and dividing by the standard deviation of the dataset. * Finally, we plot the distribution of the Z-scores using a histogram to visualize their spread.

This example showcases how the Z-score helps normalize data to a common scale, making it easier to compare different values and identify outliers in the dataset.

Sigmoid Function

$$\sigma(x) = \frac{1}{1 + e^{-x}}$$



A smooth on or off transition

Key ML Equation 4: Sigmoid Function

$$\sigma(z) = \frac{1}{1+e^{-z}}$$

- z The **input variable**, which is the linear combination of features and weights in a model. It represents the score calculated before applying the sigmoid transformation and can take any real value.
- The **sigmoid output**, which is the transformed value of z, mapped between 0 and 1. This output represents the probability that the input belongs to a particular class (e.g., 1 for positive, 0 for negative in binary classification).
- e The mathematical constant, approximately equal to 2.718, which forms the base of the natural logarithm. The exponential term e^{-z} causes the sigmoid function to asymptotically approach 0 for large negative z and 1 for large positive z.

Introduction: The sigmoid function is an activation function that outputs values between 0 and 1.

Description: It maps any input value into a range between 0 and 1, making it suitable for probability-related tasks.

Importance in ML: The sigmoid function is widely used in logistic regression and as an activation function in neural networks. It helps in modeling binary classification problems and introduces non-linearity into neural models.

4.1 What is Behind the Equation

The sigmoid function, denoted as $\sigma(x)$, is an activation function widely used in machine learning and data science. It maps any real-valued number to a value between 0 and 1, making it ideal for scenarios where probability-related interpretations are essential, especially binary classification problems.

This equation transforms an input x into an output within the range of 0 to 1, effectively normalizing the input space.

$$\sigma(x) = \frac{1}{1 + e^{-x}} = \frac{e^x}{1 + e^x} = 1 - \sigma(-x)$$
(4.1.1)

This is a usefeul identity allows for simplifications in various mathematical expressions and facilitates gradient-based optimization in machine learning models. Note that the function is differentiable, which is crucial for backpropagation in neural networks.

In many fields, particularly in artificial neural networks, the term sigmoid function usually refers to the logistic function, a specific type of S-shaped, or "sigmoid," curve. While other functions like the Gompertz curve or the ogee curve may appear similar because they share the same general S-shape, these are distinct mathematical functions with unique properties and applications.

A sigmoid function is characterized by its shape: it begins at a low value, then increases rapidly in the middle, and finally levels off at a high value. The logistic function, which is a common example, maps any real number (from negative to positive infinity) to an output between 0 and 1. This makes it especially useful when working with probabilities or values that need to be constrained within this range.

Other variations of sigmoid functions, like the hyperbolic tangent function (tanh tanh), also share the S-shaped curve but differ in their output ranges. The hyperbolic tangent function, for example, produces outputs between -1 and 1, which may be more suitable in cases where negative values are meaningful (such as for representing balanced or symmetric outcomes).

4.2 General Overview of Applications of Sigmoid Functions

• Artificial Neural Networks: Sigmoid functions are widely used as activation functions, which means they determine whether a neuron in a neural network should be "activated" (i.e., pass on its signal). In particular, the logistic function is popular because it maps values to a range between 0 and 1, simulating the all-or-nothing activation seen in

biological neurons. This behavior allows neural networks to make decisions in a gradual, continuous manner.

- Cumulative Distribution Functions in Statistics: The sigmoid function is also useful in statistics, where it can represent the cumulative distribution function (CDF) of a probability distribution. In this context, the function's smooth, continuous increase models the accumulation of probability from one extreme to another, representing how likely it is to observe values up to a certain point.
- Probabilistic Interpretation: When the output of the logistic sigmoid is interpreted as a probability (from 0 to 1), it enables the function to model binary classification problems, where there are only two possible outcomes (such as yes/no or true/false). By setting a threshold (e.g., 0.5), predictions can be made based on the probability that an input belongs to a certain class.
- Invertibility: The logistic sigmoid function is invertible, meaning it has an inverse function called the logit function. The logit function transforms probabilities (between 0 and 1) back into raw values from all real numbers, which can be useful in logistic regression and other statistical models that seek to model relationships in probability terms.

In summary, sigmoid functions, particularly the logistic function, are versatile tools in both machine learning and statistics due to their S-shape, bounded output range, and smooth transition, which makes them ideal for modeling gradual changes in various contexts.

4.3 Application to Machine Learning: Sigmoid Function in Binary Classification

In binary classification tasks, the sigmoid function is particularly valuable as it can be used to estimate the probability that a given input belongs to a certain class. Logistic regression, for example, uses the sigmoid function to model the probability of a binary outcome, with predictions being based on whether the probability exceeds a specified threshold (e.g., 0.5). By mapping outputs to a range between 0 and 1, the sigmoid function enables interpretable probability scores and smooth gradients for optimization.

4.4 How Sigmoid Works in Logistic Regression

In logistic regression, the sigmoid function serves as the "activation" that transforms the output of a linear model into a probability.

1. Linear Combination of Features: First, we calculate a linear combination of the input features. For example, in a logistic regression model with two features x_1 and x_2 , the model calculates:

$$z = w_1 x_1 + w_2 x_2 + b (4.4.1)$$

where w_1 and w_2 are weights learned from the data, and b is the bias term. This value z represents a "score" that is not yet bounded between 0 and 1.

2. Applying the Sigmoid Transformation: Next, we pass this linear combination z through the sigmoid function, which transforms it into a probability:

$$\sigma(z) = \frac{1}{1 + e^{-z}} \tag{4.4.2}$$

This output $\sigma(z)$ now represents the estimated probability that the input belongs to the positive class (e.g., "spam" or "yes").

3. Thresholding for Classification: Once we have the probability, we can decide on a threshold to classify the input. For example, if we set a threshold of 0.5, we would classify the input as belonging to the positive class if $\sigma(z) \geq 0.5$, and to the negative class otherwise. This threshold is often set based on the problem's specific requirements and the balance of classes.

4.5 Example Walkthrough

Imagine you're building a model to detect spam emails based on certain features, like the presence of certain words or the frequency of punctuation marks. Here's how the process might work step-by-step:

- 1. **Input Features**: Let's say we have two features:
 - x_1 : The frequency of the word "free."
 - x_2 : The number of exclamation marks in the email.
- 2. **Model Weights**: After training on a dataset, the model might assign the following weights:
 - $w_1 = 1.2$ (suggesting "free" is an indicator of spam)

• $w_2 = 0.8$ (suggesting a high number of exclamation marks also correlates with spam)

Let's assume the bias term b is -1.5.

3. Calculate z: For an email where $x_1 = 2$ and $x_2 = 3$ (meaning "free" appears twice and there are three exclamation marks), we calculate:

$$z = (1.2 \times 2) + (0.8 \times 3) - 1.5 = 2.4 + 2.4 - 1.5 = 3.3$$
 (4.5.1)

4. **Apply Sigmoid to Get Probability**: Now, we pass z through the sigmoid function:

$$\sigma(z) = \frac{1}{1 + e^{-3.3}} \approx 0.964 \tag{4.5.2}$$

This probability, 0.964, indicates a high likelihood that the email is spam.

5. Classification Decision: If our threshold is 0.5, we would classify this email as "spam" since 0.964 > 0.5. This probability-based approach allows us to adjust the threshold based on how conservative or lenient we want our classification to be.

4.6 Why Sigmoid is Useful for Optimization

The sigmoid function also has smooth gradients, meaning that even small changes in z lead to gradual changes in the output probability $\sigma(z)$. This smoothness is critical for optimization because it allows gradient-based algorithms, such as gradient descent, to make small, continuous updates to model parameters.

To demonstrate the smooth gradient of the sigmoid function mathematically, we can derive its derivative, which shows how the output changes with respect to changes in the input z. This derivative will show that the sigmoid function has a continuous and smooth gradient, making it suitable for gradient-based optimization methods.

4.6.1 Smooth Gradient of the Sigmoid Function

To understand why the sigmoid function has smooth gradients, let's derive its first derivative. We can calculate the derivative of $\sigma(z)$ with respect to z, which will tell us how sensitive the output $\sigma(z)$ is to small changes in z.

4.7 Deriving the Derivative of the Sigmoid Function

1. Start with the sigmoid function:

$$\sigma(z) = \frac{1}{1 + e^{-z}} \tag{4.7.1}$$

2. To find $\frac{d\sigma}{dz}$, we apply the chain rule. Let's rewrite the function in a form that will make it easier to differentiate:

$$\sigma(z) = (1 + e^{-z})^{-1} \tag{4.7.2}$$

3. Now, differentiate with respect to z:

$$\frac{d\sigma}{dz} = -(1 + e^{-z})^{-2} \cdot (-e^{-z}) \tag{4.7.3}$$

4. Simplify the expression:

$$\frac{d\sigma}{dz} = \frac{e^{-z}}{(1 + e^{-z})^2} \tag{4.7.4}$$

5. Notice that we can rewrite e^{-z} in terms of $\sigma(z)$ because:

$$\sigma(z) = \frac{1}{1 + e^{-z}} \Rightarrow e^{-z} = \frac{1 - \sigma(z)}{\sigma(z)}$$

$$(4.7.5)$$

6. Substitute this back to express $\frac{d\sigma}{dz}$ in terms of $\sigma(z)$:

$$\frac{d\sigma}{dz} = \sigma(z)(1 - \sigma(z)) \tag{4.7.6}$$

4.7.1 Interpretation of the Derivative

The derivative of the sigmoid function, $\frac{d\sigma}{dz} = \sigma(z)(1 - \sigma(z))$, shows us that:

- The gradient is always positive and smooth for all values of z.
- This derivative is largest around z = 0, where the sigmoid curve is steepest, and it becomes smaller as z moves towards positive or negative infinity, where the curve flattens out.

This smooth gradient is crucial for gradient descent, as it allows the algorithm to make gradual updates to the weights. Because the derivative never suddenly changes, the model can adjust parameters smoothly without sudden jumps, aiding in stable and effective learning.

Thus, the sigmoid's smooth gradient makes it well-suited for optimization in machine learning.

4.8 Typical Ranges or Values for the Sigmoid Function

The sigmoid function is bounded by 0 and 1. At x = 0, $\sigma(x)$ returns 0.5. As $x \to \infty$, $\sigma(x) \to 1$; and as $x \to -\infty$, $\sigma(x) \to 0$. The sigmoid function's characteristic "S"-shaped curve makes it particularly suited for gradual transitions between classes in classification tasks.

4.9 Important Mathematical Identities

Some useful identities associated with the sigmoid function include:

1. Derivative of the sigmoid:

$$\sigma'(x) = \sigma(x)(1 - \sigma(x)) \tag{4.9.1}$$

2. Relationship with the hyperbolic tangent:

$$\sigma(x) = \frac{1 + \tanh\left(\frac{x}{2}\right)}{2} \tag{4.9.2}$$

These identities assist in understanding the behavior of the sigmoid function in neural networks and other machine learning algorithms.

4.10 Comparative Functions

Several functions exhibit similar "S"-shaped or smooth transition behaviors. Here are some of them:

$$-\operatorname{erf}\left(\frac{\sqrt{\pi}}{2}x\right) \quad -\frac{x}{\sqrt{1+x^2}} \\ -\tanh(x) \quad -\frac{2}{\pi}\arctan\left(\frac{\pi}{2}x\right) \\ -\frac{2}{\pi}\operatorname{gd}\left(\frac{\pi}{2}x\right) \quad -\frac{x}{1+|x|}$$

In addition to the sigmoid function, there are several other functions that exhibit smooth, "S"-shaped curves and are used for various applications in machine learning and mathematics. Below are six of these comparative functions, along with a brief description of their unique properties.

4.10.1 Error Function:

The error function, denoted as $\operatorname{erf}(x)$, is often used in probability and statistics, particularly in relation to the normal distribution. It approximates the integral of the Gaussian function, and its shape closely resembles that of the sigmoid. The error function has applications in fields such as signal processing and heat diffusion.

$$-\operatorname{erf}\left(\frac{\sqrt{\pi}}{2}x\right) \tag{4.10.1}$$

4.10.2 Reciprocal Square Root:

This function smooths inputs similarly to the sigmoid but has a different asymptotic behavior. Unlike the sigmoid, which approaches 1 and 0, this function approaches -1 and 1 as x tends to ∞ and $-\infty$, respectively. It is sometimes used in neural networks as an alternative activation function.

$$-\frac{x}{\sqrt{1+x^2}}\tag{4.10.2}$$

4.10.3 Hyperbolic Tangent:

The hyperbolic tangent function, tanh(x), is a popular alternative to the sigmoid in neural networks because its range is from -1 to 1, allowing for more balanced gradient flows during backpropagation. It provides output symmetry, which can sometimes lead to faster training.

$$-\tanh(x) \tag{4.10.3}$$

4.10.4 Arctangent:

This function maps inputs to a range between -1 and 1 similarly to the hyperbolic tangent but has a slightly different curve. The arctangent function is sometimes used in machine learning models when smaller gradients or more gradual transitions are preferred.

$$\frac{2}{\pi}\arctan\left(\frac{\pi}{2}x\right) \tag{4.10.4}$$

4.10.5 Gudermannian Function:

The Gudermannian function, gd(x), provides a link between circular and hyperbolic functions without involving complex numbers. This function has applications in geometry and physics, particularly in mapping problems.

$$\frac{2}{\pi} \operatorname{gd}\left(\frac{\pi}{2}x\right) \tag{4.10.5}$$

4.10.6 Inverse Absolute:

This function smoothly scales inputs, but its range only extends from -1 to 1. Its asymptotic properties are particularly useful when a smoother transition around zero is preferred.

$$\frac{x}{1+|x|}\tag{4.10.6}$$

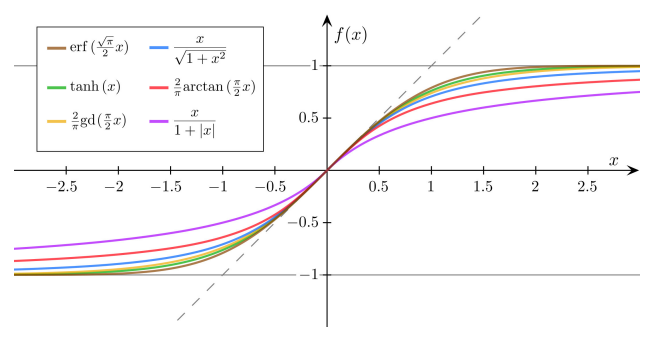


Figure 4.1: Summary of Alternate Sigmoid Like Curves

Note: All functions are normalized in such a way that their slope at the origin is 1

Correlation

Correlation =
$$\frac{\text{Cov}(X,Y)}{\text{Std}(X)\cdot\text{Std}(Y)}$$



 \boldsymbol{x}_i and \boldsymbol{y}_i are linearly related

Key ML Equation 5: Correlation

$$Correlation = \frac{Cov(X, Y)}{Std(X) \cdot Std(Y)}$$
 (5.0.1)

(5.0.2)

$$r = \frac{\sum (x_i - \bar{x})(y_i - \bar{y})}{\sqrt{\sum (x_i - \bar{x})^2 \sum (y_i - \bar{y})^2}}$$
 (5.0.3)

- Cov(X, Y) The **covariance** between vectors X and Y, which measures the degree to which the two variables (or vectors) vary together.
- Std(V) The standard deviation of vector V, which measures the spread of the elements of Y around each mean.
- \bar{x} The **mean of variable** X, calculated as $\frac{1}{n} \sum x_i$, where n is the number of data points.
- \bar{y} The **mean of variable** Y, calculated as $\frac{1}{n} \sum y_i$, where n is the number of data points.
- x_i The **individual data point** in the variable X, where i indexes each observation in the dataset.
- y_i The **individual data point** in the variable Y, where i indexes each observation in the dataset.

Introduction: Correlation measures the strength and direction of the linear relationship between two variables.

Description: It ranges from -1 to 1, where values close to 1 or -1 indicate a strong relationship, and values close to 0 indicate no relationship.

Importance in ML: Correlation analysis helps in feature selection by identifying which features are related to the target variable. Strongly correlated features can be used for better predictions, whereas uncorrelated features can be removed to simplify models.

5.1 What is Behind the Equation

Correlation is a statistical measure that describes the strength and direction of a relationship between two variables. The most common measure of correlation is Pearson's correlation coefficient, denoted as r, which ranges from -1 to 1. A positive value indicates a direct relationship, while a negative value indicates an inverse relationship. A value of 0 implies no linear relationship.

The equation for Pearson's correlation r coefficient 5.0.3 can be compared to the normalized version 5.1.1:

$$r_{\mu} \frac{\sum (x_i y_i)}{\sqrt{\sum x_i^2 \sum y_i^2}} \tag{5.1.1}$$

Where: - x_i and y_i are individual data points of variables x and y, - \bar{x} and \bar{y} are the means of variables x and y, respectively.

- Variables: The two variables, x and y, whose relationship we are measuring. These could be anything from economic indicators to user behaviors.
- Data Points: Each pair of values (x_i, y_i) represents an observation.
- Means: The mean values of x and y are used to standardize the data by subtracting them from each data point, which helps in normalizing the relationship.
- x_i and y_i are individual data points for the two variables X and Y.
- \bar{x} and \bar{y} are the means (averages) of the respective variables.
- **Numerator**: The numerator is the **covariance** between *X* and *Y*, which measures how the two variables vary together.
- **Denominator**: The denominator is the product of the **standard deviations** of *X* and *Y*, which normalizes the covariance and ensures that the correlation coefficient lies between -1 and 1.

The Pearson correlation measures the strength and direction of the **linear relationship** between two variables. This is the most commonly used form of correlation.

5.1.0.1 Use case:

This equation is used when you want to understand the **linear relationship** between two continuous variables and is the formula used by the **cor()** function in R with the default method (method = "pearson").

5.1.0.2 Difference in Interpretation:

- The pearson equation 5.0.3 measures how deviations from the mean of each variable are related, i.e., it normalizes by the standard deviations of each variable.
- The second equation 5.1.1 compares the raw products of the variables, but it does **not center** the data by subtracting the means. This makes the second equation more sensitive to the scales and magnitudes of the variables, which is why it's often less preferred for calculating correlation in its standard sense.

5.1.1 Key Differences:

1. Normalization:

- The first equation involves centering the data by subtracting the means, which is important to account for the relationship of the variables in terms of deviations from their respective averages.
- The second equation does not involve centering the data. It uses the raw values of x_i and y_i , making it sensitive to the overall scale and magnitude of the data.

2. Covariance vs Raw Product:

- The first equation is essentially computing a normalized **covariance** (divided by the product of the standard deviations of *X* and *Y*).
- The second equation uses the **raw product** of the variables, which lacks the standardization and may lead to different numerical results.

3. Interpretation:

- The first equation (Pearson's correlation) measures the strength of a linear relationship, normalized by the variance in both variables.
- The second equation is more general but is not commonly used as a standard
 for measuring correlation. It may appear in specific contexts, such as cosine
 similarity when the data is already standardized or when normalization is not
 necessary.

5.1.2 Conclusion:

- The first equation ?? is the Pearson correlation coefficient, the standard method for measuring linear relationships, normalized by the standard deviations of both variables.
- The second equation 5.1.1 is a raw form that compares the products of the variables

and is **not typically used** for Pearson's correlation, as it lacks normalization. It could be seen as a raw form of a **cosine similarity** (for vector comparison) or some other variant of correlation, but it's not generally used for typical correlation measurements.

5.2 5.3 Historical Context

The concept of correlation was first developed by Sir Francis Galton in the 19th century. Galton's work laid the foundation for the field of regression analysis and correlational statistics. The Pearson correlation, developed by Karl Pearson in 1896, is the most widely used form of correlation in modern statistics.

5.3 5.4 Understanding the Notation

The formula for Pearson's correlation coefficient involves the concept of covariance, which measures how much two variables change together. The denominator in the formula standardizes the covariance by dividing by the product of the standard deviations of x and y.

- Covariance: Measures the degree to which two variables move together.
- Standard Deviation: Measures the spread of a variable around its mean.

5.4 5.5 Applications in Machine Learning

In machine learning, correlation is used to: - Identify relationships between input features and target variables, - Check for multicollinearity, where highly correlated features might skew results, - Perform feature selection, as correlated features may provide redundant information.

For instance, if two features are highly correlated, one may be removed from the model to simplify it without losing predictive power.

5.5 5.6 Correlation in R

To compute the correlation coefficient in R, you can use the built-in cor() function:

```
# Example R code to compute Pearson's correlation coefficient
x <- c(1, 2, 3, 4, 5)
y <- c(5, 4, 3, 2, 1)
correlation <- cor(x, y)
# print{correlation}
cat(x)
## 1 2 3 4 5
cat(y)</pre>
```

5 4 3 2 1

cat(correlation)

-1

Cosine Similarity

$$\text{similarity} = \frac{A \cdot B}{\|A\| \|B\|}$$

Introduction: Cosine similarity measures the cosine of the angle between two vectors in a multi-dimensional space.

Description: It is commonly used to determine how similar two documents are, regardless of their size, by comparing their word vectors.

Importance in ML: Cosine similarity is crucial in text mining and information retrieval applications. It is often used in clustering and classification algorithms for textual data, where measuring the similarity between vectors is needed.

Naive Bayes

$$P(y|x_1,\ldots,x_n) = \frac{P(y)\prod_{i=1}^n P(x_i|y)}{P(x_1,\ldots,x_n)}$$

Introduction: Naive Bayes is a probabilistic classifier based on Bayes' theorem with strong independence assumptions.

Description: It assumes that the presence of a particular feature in a class is unrelated to the presence of any other feature, which simplifies computation.

Importance in ML: Naive Bayes is used for classification tasks, especially in text classification and spam detection. Its simplicity and efficiency make it a popular choice for high-dimensional datasets.

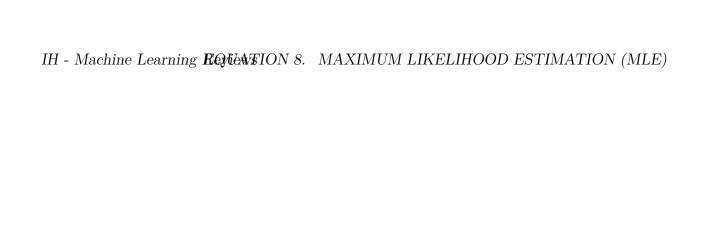
Maximum Likelihood Estimation (MLE)

$$\arg\max_{\theta} \prod_{i=1}^{n} P(x_i|\theta)$$

Introduction: MLE is a method used to estimate the parameters of a statistical model.

Description: It finds the parameter values that maximize the likelihood of making the observations given the model.

Importance in ML: MLE is used to train many machine learning models, including logistic regression and Gaussian mixture models. It provides a way to fit models to data and make statistical inferences.



Ordinary Least Squares (OLS)

$$\hat{\beta} = (X^T X)^{-1} X^T y$$

Introduction: OLS is a method for estimating the unknown parameters in a linear regression model.

Description: It minimizes the sum of squared residuals between the observed and predicted values.

Importance in ML: OLS is a fundamental technique for regression analysis. It helps determine the best-fit line for the data, making it useful for predictive modeling.

F1 Score

$$F_1 = \frac{2 \cdot P \cdot R}{P + R}$$

Introduction: The F1 score is a measure of a test's accuracy that considers both precision (P) and recall (R).

Description: It is the harmonic mean of precision and recall, providing a single metric to evaluate model performance.

Importance in ML: The F1 score is particularly useful for imbalanced datasets where the distribution of classes is uneven, providing a balanced measure of model effectiveness.

ReLU (Rectified Linear Unit)

$$ReLU(x) = max(0, x)$$

Introduction: ReLU is an activation function used in neural networks.

Description: It outputs the input directly if positive; otherwise, it outputs zero.

Importance in ML: ReLU is crucial for deep learning as it helps mitigate the vanishing gradient problem, allowing models to learn faster and perform better.

Softmax Function

$$P(y=j|z) = \frac{e^{zw_j}}{\sum_{k=1}^K e^{zw_k}}$$

Introduction: Softmax is an activation function that outputs a probability distribution over multiple classes.

Description: It converts raw scores into probabilities, which sum up to one, making it suitable for multi-class classification.

Importance in ML: Softmax is commonly used in the output layer of neural networks for classification problems involving multiple classes, making it essential for multi-class models.

R-squared (R^2) Score

$$R^2 = 1 - \frac{\sum_{i=1}^n (y_i - \hat{y}_i)^2}{\sum_{i=1}^n (y_i - \bar{y})^2}$$

Introduction: R-squared is a statistical measure that represents the proportion of variance in the dependent variable explained by the independent variables.

Description: It is a goodness-of-fit measure that ranges from 0 to 1, with values closer to 1 indicating a better fit.

Importance in ML: R-squared is used to evaluate the performance of regression models. It provides insights into how well the model explains the variance in the target variable.

Mean Squared Error (MSE)

$$MSE = \frac{1}{n} \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$

Introduction: Mean Squared Error is a common loss function used to measure the average squared difference between predicted and actual values.

Description: It calculates the squared difference for each observation and then averages them to provide a metric of model accuracy.

Importance in ML: MSE is used to evaluate the performance of regression models. Lower MSE values indicate a better fit, making it essential for identifying the optimal model.

Mean Squared Error with L2 Regularization (MSE + L2 Reg)

$$\text{MSE}_{\text{regularized}} = \frac{1}{n} \sum_{i=1}^{n} (y_i - \hat{y}_i)^2 + \lambda \sum_{j=1}^{p} \beta_j^2$$

Introduction: Regularized MSE adds a penalty to the loss function to prevent overfitting by discouraging overly complex models.

Description: The regularization term $\lambda \sum \beta_j^2$ helps keep model parameters small, which leads to simpler models.

Importance in ML: L2 regularization is key in reducing overfitting by controlling the complexity of the model, ensuring that the model generalizes well to new data.

Eigenvectors and Eigenvalues

 $Av = \lambda v$

Introduction: Eigenvectors and eigenvalues are fundamental concepts in linear algebra used to understand the structure of matrices.

Description: They represent the directions along which linear transformations act by only scaling the vectors, without changing their direction.

Importance in ML: Eigenvectors and eigenvalues are used in dimensionality reduction techniques like PCA (Principal Component Analysis), which helps reduce the number of features while preserving essential information.

Entropy

$$\text{Entropy} = -\sum_i p_i \log_2(p_i)$$

Introduction: Entropy is a measure of uncertainty or randomness in a dataset.

Description: It quantifies the impurity in a dataset, making it a key concept in information theory and decision trees.

Importance in ML: Entropy is used in decision tree algorithms to decide the best split at each node by measuring the purity of a dataset. Lower entropy indicates a more homogenous group of samples.

K-Means Clustering

$$\arg\min_{S} \sum_{k=1}^K \sum_{x \in S_k} \|x - \mu_k\|^2$$

Introduction: K-Means is an unsupervised learning algorithm used for clustering data into K distinct groups.

Description: It minimizes the sum of squared distances between data points and the centroid of their assigned cluster.

Importance in ML: K-Means is a fundamental clustering technique used in exploratory data analysis and segmentation tasks. It helps discover patterns and relationships in unlabeled data.

Kullback-Leibler (KL) Divergence

$$D_{KL}(P||Q) = \sum_{x \in \mathcal{X}} P(x) \log \frac{P(x)}{Q(x)}$$

Introduction: KL Divergence is a measure of how one probability distribution diverges from a second, reference probability distribution.

Description: It is commonly used to measure the difference between two probability distributions.

Importance in ML: KL Divergence is used in machine learning for loss functions in models like variational autoencoders. It quantifies how well the learned distribution approximates the true data distribution.

IH - Machine Learning ReviewUATION 19.	$KULLBACK\text{-}LEIBLER\ (KL)\ DIVERGENCE$

Log Loss

$$-\frac{1}{N} \sum_{i=1}^{N} \left(y_i \log(\hat{y}_i) + (1-y_i) \log(1-\hat{y}_i) \right)$$

Introduction: Log Loss, or logistic loss, is a loss function used for binary classification tasks.

Description: It measures the performance of a classification model whose output is a probability value between 0 and 1.

Importance in ML: Log Loss is crucial in evaluating classification models where the output is a probability. Lower log loss indicates a more accurate model, especially for probabilistic predictions.

Support Vector Machine (SVM) Objective

$$\min_{w,b} \frac{1}{2} \|w\|^2 + C \sum_{i=1}^n \max(0, 1 - y_i(w \cdot x_i - b))$$

Introduction: SVM is a supervised learning model used for classification and regression tasks.

Description: It finds the hyperplane that best separates different classes by maximizing the margin between them.

Importance in ML: SVMs are effective for high-dimensional spaces and are used in classification problems where the decision boundary is non-linear.

Linear Regression

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \dots + \beta_n x_n + \epsilon$$

Introduction: Linear regression is a simple and widely used method for predictive analysis.

Description: It models the relationship between a dependent variable and one or more independent variables by fitting a linear equation to observed data.

Importance in ML: Linear regression is fundamental for understanding relationships between variables and is used in predictive modeling to estimate outcomes.

Singular Value Decomposition (SVD)

$$A = U\Sigma V^T$$

Introduction: SVD is a matrix factorization technique used in linear algebra.

Description: It decomposes a matrix into three other matrices, revealing the intrinsic structure of the data.

Importance in ML: SVD is used in dimensionality reduction, data compression, and noise reduction. It is also a core algorithm behind recommendation systems.

Lagrange Multiplier

$$\max f(x) \; ; \; g(x) = 0$$

$$L(x,\lambda) = f(x) - \lambda * g(x)$$

Introduction: The Lagrange multiplier is a method for finding the local maxima and minima of a function subject to equality constraints.

Description: It introduces a new variable, λ , which helps in optimizing a function while considering the constraint.

Importance in ML: Lagrange multipliers are used in optimization problems with constraints, such as training machine learning models with regularization terms.

The Human Equation

HUMAN

Introduction:

Description:

Importance in ML:

Epilogue - Guidance and Test Materials

Applying Musk Rules to Writing Your Book on the "25 Key Equations in Machine Learning"

1. Always Question the Requirements

- Book Scope and Audience: Reevaluate the core purpose and audience of the book. Who is your ideal reader? Do you really need to cover 25 equations, or would a different number be more impactful? Is there any unnecessary content that isn't truly essential to your goal of effective teaching and reference?
- **Key Questions to Ask:** Do each of the sections contribute to the core learning objectives? Could some be combined, shortened, or presented differently to be clearer and more engaging?

2. Try Very Hard to Delete Parts or Processes

- **Delete Unnecessary Content:** Go through each chapter and decide if every subsection or detail is truly required. Are there parts of the explanations, historical contexts, or code examples that could be condensed or cut without losing value?
- Focus on Core Concepts: Sometimes, less is more. Identify and focus on the key insights of each equation. For example, rather than going deep into every historical development of the normal distribution, focus on how the normal distribution is applied directly in machine learning.

3. Simplify or Optimize, But Not Too Early

• Simplification After Completion: In the initial phase, write freely to capture all your thoughts and ideas. Don't worry too much about the length or complexity. Once you have a draft, then simplify—remove verbose explanations, simplify complex code

snippets, or summarize complex derivations to keep it engaging.

• Iteration in Drafts: Write each chapter in an exploratory way first, then optimize the language and reduce complexity in subsequent drafts.

4. Move Faster

- Accelerate Writing with LLMs and Tools: Use tools like large language models (LLMs) to help generate content, summarize, and write initial drafts faster. You've already indicated using Mathpix, LaTeX, and R Markdown—lean into these tools to keep your process fast and flexible.
- Set Short-Term Goals: Use milestones to maintain momentum. Instead of working on all 25 chapters at once, work on drafting 5 chapters in a week. Use techniques like "pomodoro" to keep the pace fast without burnout.

5. Finally: Automate

- Automate Repetitive Tasks Last: Once content is ready, automate formatting and typesetting. Given that you are using R Markdown and LaTeX, continue leveraging those tools for repetitive typesetting tasks, but do this after you've streamlined the core content.
- Automation for Consistency: Automate consistency checks for formulas and code. This could be done with scripts that verify equation formatting, syntax checking for code snippets, or tools that flag unformatted variables.

Specific Strategies to Speed Up the Book Creation:

1. Outline Consolidation:

Review your current outline and condense or merge sections that overlap. Consider
if all 25 equations deserve equal weight. You could provide detailed explanations
for 10-15 equations while giving more general summaries for the rest if they are
less foundational.

2. Minimize and Prioritize:

• Focus on the most impactful content. Given the target audience, ask yourself: do they need a historical deep dive, or is the practical application what matters most? Emphasize applications, examples, and visuals over long theoretical discussions unless absolutely necessary.

3. Visuals and Examples:

• To make learning engaging, visuals and practical examples are key. Automate the creation of code snippets and figures that are repetitive. Tools like R and Python

scripts can automatically generate the visuals needed for each equation, saving considerable time.

4. Leverage Existing Content:

• Since you are creating this as a reference book, consider using existing open-source implementations of the concepts as examples, rather than creating everything from scratch. This will save time while still providing value.

5. Peer Review and Feedback Early:

• Incorporate early feedback to streamline revisions. Share your draft with a small group of readers and ask for their feedback on clarity and structure. This will help you catch issues early, saving time during the editing phase.

6. Use a Versioned Approach:

• Set smaller, achievable versions (like v0.3, v0.4, etc.) where each iteration refines content. For example, v0.3 might be the initial full draft, v0.4 could focus on reducing complexity, and v0.5 could involve formatting and visual consistency. This will help you move in structured steps rather than getting bogged down in perfecting each detail upfront.

Moving Faster, in Practice:

- Sprint Write Sections: Work on chapters in sprints. Set a specific time limit for each chapter, such as three days per equation. The goal is to create momentum and avoid stagnation.
- Lean on a Support System: Delegate part of the process. If possible, outsource parts like copy-editing, formatting, or even generating diagrams. The faster you move past the basic draft stage, the better you can refine and make the book truly impactful.

Conclusion:

The Musk Rules encourage challenging the purpose of everything you're including, focusing on essential content, and using tools and automation wisely, but only after the core content is strong. By questioning, deleting, simplifying, moving fast, and automating thoughtfully, you can accelerate the process of creating your book without compromising quality. Focus on the main principles that will make your book a genuinely valuable reference—clear, effective, and engaging explanations, visualizations, and examples—while avoiding unnecessary details that slow you down.

How do these recommendations feel in the context of your goals for this book? Would you like help refining a specific section or strategy for moving faster?

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25.1 TEMP _ R code - Python code - test area

25.2 testing Python Integration (temp section - to remove later)

Create a variable ${\tt x}$ in the Python session:

$$x = [1, 2, 3]$$

Access the Python variable x in an R code chunk:

py\$x

[1] 1 2 3

Create a new variable y in the Python session using R, and pass a data frame to y:

py\$y <- head(cars)</pre>

Print the variable y in Python:

print(y)

{'speed': [4.0, 4.0, 7.0, 7.0, 8.0, 9.0], 'dist': [2.0, 10.0, 4.0, 22.0, 16.0, 10.

25.3 Define Numbers in R

Let's define the numbers we will use in both R and Python:

```
# Define a vector of numbers in R
define_numbers <- c(1, 2, 3, 4, 5)</pre>
```

25.4 R Code Block

This is a simple R code block that calculates the sum of 5 numbers:

```
# R code to sum 5 numbers
r_sum <- sum(define_numbers)
print(paste("The sum of the numbers in R is:", r_sum))</pre>
```

[1] "The sum of the numbers in R is: 15"

25.5 Python Code To Pass *

```
py$n2 <- define_numbers</pre>
```

25.6 Python Code Block

This is a simple Python code block that calculates the sum of 5 numbers:

```
# Python code to calculate the sum of numbers defined in R

# Define a list of numbers
numbers = [1, 2, 3, 4, 5]

# Retrieve the numbers passed from R using reticulate's 'py' object
total_sum = sum(numbers)
total_sum2 = sum(n2)

# Print the result
print(f"The sum of the numbers 'total sum' is: {total_sum}")
```

The sum of the numbers 'total sum' is: 15

```
\label{lem:print(f"The sum of the numbers 'n2' is: {total\_sum2}")} \\
```

The sum of the numbers 'n2' is: 15.0

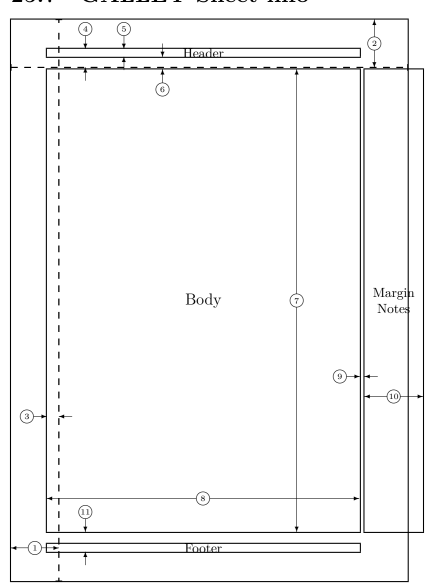
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Galley Sheet

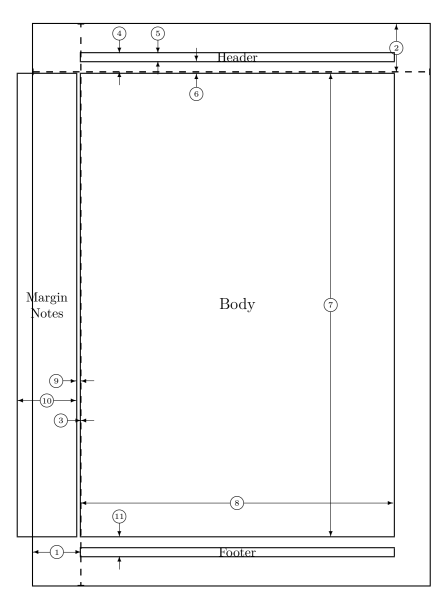
25.7 GALLEY Sheet info



- 1 one inch + \hoffset
- 3 \oddsidemargin = −18pt
- 5 \headheight = 12pt
- 7 \textheight = 696pt
- 9 \marginparsep = 7pt
- 11 \footskip = 30pt
 \hoffset = 0pt
 \paperwidth = 597pt
- 2 one inch + \voffset
- 4 \topmargin = -28pt
- 6 \headsep = 19pt
- 8 \textwidth = 471pt

10 \marginparwidth = 88pt

- \marginparpush = 7pt (not shown)
 - \voffset = Opt
 - \paperheight = 845pt



- 1 one inch + \hoffset
- 3 \evensidemargin = Opt
- 5 \headheight = 12pt
- 7 \textheight = 696pt
- 9 \marginparsep = 7pt
- 11 \footskip = 30pt
 \hoffset = 0pt
 \paperwidth = 597pt
- 2 one inch + \voffset
- 4 \forall topmargin = -28pt
- 6 \headsep = 19pt
- $8 \quad \text{textwidth} = 471pt$
- 10 \marginparwidth = 88pt
 \marginparpush = 7pt (not shown)
 \voffset = 0pt
 \paperheight = 845pt

25.8 GALLEY Sheet info - ends

References