

# Week 0: Introduction to Machine Learning & AI

## Foundations, Algorithms, and Modern Applications

Machine Learning for Smarter Innovation

BSc-Level Course Series

September 28, 2025

1 Machine Learning Foundations

2 Supervised Learning Methods

3 Unsupervised Learning Methods

# Part 1: Machine Learning Foundations

## Theory, Definitions, and Core Concepts

## Tom Mitchell's Definition (1997)

A computer program learns from:

- **Experience  $E$**  with respect to
- **Task  $T$**  and
- **Performance measure  $P$**

if its performance at task  $T$ , as measured by  $P$ , improves with experience  $E$ .

### Example:

E: Email database

T: Classify spam vs non-spam

P: Classification accuracy

## Mathematical View

Learn function  $f : \mathcal{X} \rightarrow \mathcal{Y}$

Given training set:

$$\mathcal{D} = \{(x_i, y_i)\}_{i=1}^n$$

Find:

$$\hat{f} =_{f \in \mathcal{F}} \sum_{i=1}^n L(y_i, f(x_i)) + \lambda R(f)$$

where:

- $L$ : Loss function
- $R$ : Regularization term
- $\lambda$ : Regularization strength

# Three Paradigms of Machine Learning

## Supervised



$$\{(x_i, y_i)\}_{i=1}^n \rightarrow \hat{f}$$

### Applications:

- Email spam detection
- Medical diagnosis
- Stock price prediction
- Image recognition

### Key Algorithms:

- Linear Regression
- Random Forest
- Neural Networks

## Unsupervised



$$\{x_i\}_{i=1}^n \rightarrow \text{Structure}$$

### Applications:

- Customer segmentation
- Anomaly detection
- Data compression
- Market basket analysis

### Key Algorithms:

- K-means clustering
- PCA
- Autoencoders

## Reinforcement



$$(s_t, a_t, r_t, s_{t+1}) \rightarrow \pi^*$$

### Applications:

- Game playing (Chess, Go)
- Autonomous vehicles
- Robotics control
- Resource allocation

### Key Algorithms:

- Q-Learning
- Policy Gradients
- Actor-Critic

# The Machine Learning Pipeline

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## Mathematical Framework

For any learning algorithm, the expected error can be decomposed as:

$$\text{Error} = \text{Bias}^2 + \text{Variance} + \text{Noise}$$

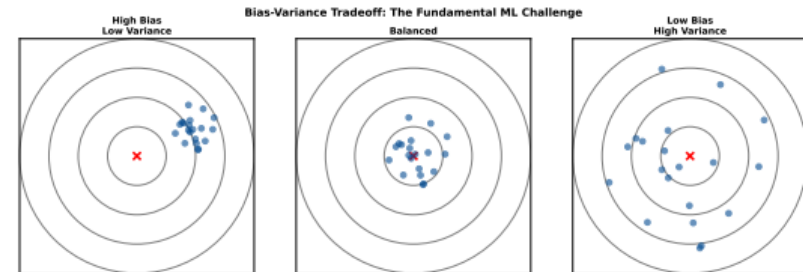
**Bias:** Error from oversimplifying assumptions

$$\text{Bias}[\hat{f}(x)] = E[\hat{f}(x)] - f(x)$$

**Variance:** Error from sensitivity to training data

$$\text{Var}[\hat{f}(x)] = E[(\hat{f}(x) - E[\hat{f}(x)])^2]$$

**Key Insight:** There's a fundamental tradeoff between bias and variance



### Model Complexity Examples:

- **High Bias:** Linear models on nonlinear data
- **Balanced:** Regularized models
- **High Variance:** Deep trees, k-NN with small k

## Traditional Programming

### Process:

- Write explicit rules
- Code logic step by step
- Handle edge cases manually
- Deterministic outputs

### Example: Email Classification

- IF contains “FREE” AND “LIMITED TIME”
- THEN classify as spam
- Requires manual rule updates

### Limitations:

- Rules become complex
- Hard to handle exceptions
- Doesn't adapt to new patterns

## Machine Learning

### Process:

- Provide example data
- Algorithm learns patterns
- Generalizes to new cases
- Probabilistic outputs

### Example: Email Classification

- Train on 10,000 labeled emails
- Learn complex word patterns
- Automatically adapts to new spam

### Advantages:

- Handles complex patterns
- Adapts to new data
- Discovers hidden relationships

## Why Split Data?

### Training Set (60%):

- Used to fit model parameters
- Algorithm learns from this data
- Larger is generally better

### Validation Set (20%):

- Used for hyperparameter tuning
- Model selection and comparison
- Prevents overfitting to training data

### Test Set (20%):

- Final unbiased evaluation
- Never seen during development
- Estimates real-world performance

charts/data\_splitting.pdf

## Classification Metrics

Accuracy:

$$\text{Accuracy} = \frac{\text{Correct Predictions}}{\text{Total Predictions}}$$

Precision:

$$\text{Precision} = \frac{\text{True Positives}}{\text{True Positives} + \text{False Positives}}$$

Recall (Sensitivity):

$$\text{Recall} = \frac{\text{True Positives}}{\text{True Positives} + \text{False Negatives}}$$

F1-Score:

$$F1 = 2 \cdot \frac{\text{Precision} \times \text{Recall}}{\text{Precision} + \text{Recall}}$$

## Regression Metrics

Mean Squared Error:

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

Root Mean Squared Error:

$$RMSE = \sqrt{MSE}$$

Mean Absolute Error:

$$MAE = \frac{1}{n} \sum_{i=1}^n |y_i - \hat{y}_i|$$

R-squared:

$$R^2 = 1 - \frac{SS_{res}}{SS_{tot}}$$

## Part 2: Supervised Learning Methods

### Prediction and Classification Algorithms

## Linear Regression Family

Ordinary Least Squares:

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

$$\min_{\beta} \|y - X\beta\|_2^2$$

Ridge Regression (L2):

$$\hat{\beta}_{ridge} = (X^T X + \lambda I)^{-1} X^T y$$

$$\min_{\beta} \|y - X\beta\|_2^2 + \lambda \|\beta\|_2^2$$

LASSO (L1):

$$\min_{\beta} \|y - X\beta\|_2^2 + \lambda \|\beta\|_1$$

No closed form - use coordinate descent

charts/linear\_regression\_comparison.pdf

### Applications:

House price prediction

## Mathematical Framework

Logistic Function:

$$p(y = 1|x) = \frac{1}{1 + e^{-(\beta_0 + \beta^T x)}}$$

Odds Ratio:

$$\frac{p}{1-p} = e^{\beta_0 + \beta^T x}$$

Log-Likelihood:

$$\ell(\beta) = \sum_{i=1}^n [y_i \log p_i + (1 - y_i) \log(1 - p_i)]$$

No closed form solution

- Use gradient descent
- Newton-Raphson method
- Iteratively reweighted least squares

charts/logistic\_regression.pdf

Decision Boundary:

$$\beta_0 + \beta^T x = 0$$

## Optimization Problem

Primal Form:

$$\min_{w,b} \frac{1}{2} \|w\|^2$$

$$\text{s.t. } y_i(w^T x_i + b) \geq 1, \forall i$$

Dual Form:

$$\max_{\alpha} \sum_i \alpha_i - \frac{1}{2} \sum_{i,j} \alpha_i \alpha_j y_i y_j x_i^T x_j$$

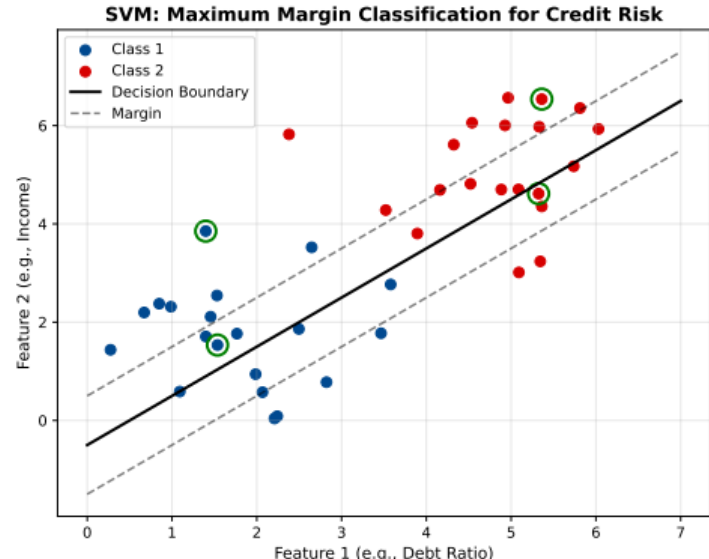
$$\text{s.t. } \alpha_i \geq 0, \sum_i \alpha_i y_i = 0$$

Kernel Trick:

$$K(x_i, x_j) = \phi(x_i)^T \phi(x_j)$$

Common kernels:

- RBF:  $K(x, z) = e^{-\gamma \|x-z\|^2}$



### Key Concepts:

- Support Vectors:** Data points on margin
- Maximum Margin:** Optimal separating hyperplane
- Kernel Trick:** Nonlinear classification

### Advantages:

• Works well in high dimensions

## Tree Construction

Splitting Criterion:

*Gini Impurity:*

$$G = \sum_{k=1}^K p_k(1 - p_k)$$

*Information Gain:*

$$IG = H(\text{parent}) - \sum_j \frac{n_j}{n} H(\text{child}_j)$$

*Entropy:*

$$H = - \sum_{k=1}^K p_k \log_2 p_k$$

**CART Algorithm:**

- ① Find best split across all features
- ② Partition data based on split
- ③ Repeat until stopping rule

charts/decision\_tree.pdf

**Advantages:**

- Highly interpretable
- Handles mixed data types

## Ensemble Method

### Bootstrap Aggregating (Bagging):

- ① Draw  $B$  bootstrap samples
- ② Train tree on each sample
- ③ Average predictions (regression)
- ④ Vote on class (classification)

### Random Feature Selection:

- At each split, randomly select  $m$  features
- Typically  $m = \sqrt{p}$  for classification
- Typically  $m = p/3$  for regression

### Final Prediction:

$$\hat{y} = \frac{1}{B} \sum_{b=1}^B T_b(x)$$

**Key insight:** Averaging reduces variance while maintaining low bias

charts/random\_forest.pdf

### Advantages:

- Reduces overfitting
- Handles missing values

## Boosting Algorithm

Sequential Model Building:

$$F_m(x) = F_{m-1}(x) + \gamma_m h_m(x)$$

where  $h_m$  is trained on residuals:

$$r_{im} = -\frac{\partial L(y_i, F_{m-1}(x_i))}{\partial F_{m-1}(x_i)}$$

XGBoost Objective:

$$\mathcal{L} = \sum_i l(y_i, \hat{y}_i) + \sum_k \Omega(f_k)$$

$$\Omega(f) = \gamma T + \frac{1}{2} \lambda ||w||^2$$

Key Features:

- Regularization prevents overfitting
- Second-order derivatives
- Handles missing values

Popular Implementations:

- **XGBoost:** Extreme Gradient Boosting
- **LightGBM:** Fast gradient boosting

charts/gradient\_boosting.pdf

## Non-parametric Method

### Algorithm:

- ① Store all training data
- ② For new point, find k nearest neighbors
- ③ Classification: majority vote
- ④ Regression: average target values

### Distance Metrics:

- Euclidean:  $d(x, z) = \sqrt{\sum_i (x_i - z_i)^2}$
- Manhattan:  $d(x, z) = \sum_i |x_i - z_i|$
- Minkowski:  $d(x, z) = (\sum_i |x_i - z_i|^p)^{1/p}$

### Choosing k:

- Small k: Low bias, high variance
- Large k: High bias, low variance
- Use cross-validation to select

charts/knn\_classification.pdf

### Advantages:

- Simple to understand and implement
- No assumptions about data distribution

# Supervised Learning: Algorithm Comparison

charts/algorithm\_comparison.pdf

## Part 3: Unsupervised Learning Methods

### Discovering Hidden Structure in Data

## Algorithm

### Objective Function:

$$J = \sum_{i=1}^n \sum_{k=1}^K w_{ik} \|x_i - \mu_k\|^2$$

### K-means Algorithm:

- ① Initialize K cluster centers randomly
- ② Assign each point to nearest center
- ③ Update centers to mean of assigned points
- ④ Repeat until convergence

### Update Rules:

$$w_{ik} = \begin{cases} 1 & \text{if } k = j \\ 0 & \text{otherwise} \end{cases}$$

$$\mu_k = \frac{\sum_{i=1}^n w_{ik} x_i}{\sum_{i=1}^n w_{ik}}$$

charts/kmeans\_clustering.pdf

### Choosing K:

- **Elbow Method:** Plot within-cluster sum of squares
- **Silhouette Analysis:** Measure cluster separation

## Agglomerative Approach

### Algorithm:

- ① Start with each point as its own cluster
- ② Merge closest pair of clusters
- ③ Repeat until single cluster remains
- ④ Cut dendrogram at desired level

### Linkage Criteria:

- **Single:**  $\min(d(a, b))$  where  $a \in A, b \in B$
- **Complete:**  $\max(d(a, b))$  where  $a \in A, b \in B$
- **Average:**  $\frac{1}{|A||B|} \sum_{a \in A} \sum_{b \in B} d(a, b)$
- **Ward:** Minimize within-cluster variance

**Time Complexity:**  $O(n^3)$  for naive implementation

charts/hierarchical\_clustering.pdf

### Advantages:

- No need to specify number of clusters
- Produces hierarchy of clusters

## Density-Based Approach

### Key Concepts:

- **Core Point:**  $\geq \text{minPts}$  neighbors within  $\epsilon$
- **Border Point:** In neighborhood of core point
- **Noise Point:** Neither core nor border

### Algorithm:

- ① For each unvisited point
- ② If core point, start new cluster
- ③ Add all density-reachable points
- ④ Mark non-core points as noise

### Parameters:

- $\epsilon$ : Neighborhood radius
- minPts: Minimum points for core

$$\text{Density} = \frac{\text{Points in } \epsilon\text{-neighborhood}}{|\epsilon\text{-neighborhood}|}$$

charts/dbSCAN\_clustering.pdf

### Advantages:

- Finds arbitrary-shaped clusters
- Automatically determines cluster count

## Mathematical Framework

**Objective:** Find directions of maximum variance

**Covariance Matrix:**

$$C = \frac{1}{n-1} X^T X$$

**Eigendecomposition:**

$$C = V \Lambda V^T$$

where  $V$  contains eigenvectors (principal components) and  $\Lambda$  contains eigenvalues.

**Dimensionality Reduction:**

$$Z = XW$$

where  $W$  contains the first  $k$  principal components.

**Variance Explained:**

$$\text{Explained Variance} = \frac{\sum_{i=1}^k \lambda_i}{\sum_{i=1}^p \lambda_i}$$

charts/pca\_analysis.pdf

### Steps:

- ① Standardize the data
- ② Compute covariance matrix

## Architecture

Encoder:

$$z = f(Wx + b)$$

Decoder:

$$\hat{x} = g(W'z + b')$$

Objective:

$$\min_{W, W'} ||x - \hat{x}||^2$$

Types of Autoencoders:

- **Vanilla:** Basic encoder-decoder
- **Denoising:** Add noise to input
- **Sparse:** Encourage sparse representations
- **Variational:** Probabilistic latent space

Bottleneck Layer: Forces compression and learning of important features

charts/autoencoder\_architecture.pdf

Advantages over PCA:

- Nonlinear transformations
- Better reconstruction for complex data

## t-SNE

### t-Distributed Stochastic Neighbor Embedding

High-dimensional similarities:

$$p_{j|i} = \frac{\exp(-\|x_i - x_j\|^2 / 2\sigma_i^2)}{\sum_{k \neq i} \exp(-\|x_i - x_k\|^2 / 2\sigma_i^2)}$$

Low-dimensional similarities:

$$q_{ij} = \frac{(1 + \|y_i - y_j\|^2)^{-1}}{\sum_{k \neq l} (1 + \|y_k - y_l\|^2)^{-1}}$$

Objective: Minimize KL divergence

$$C = \sum_i KL(P_i || Q_i)$$

Key Features:

- Preserves local structure
- Heavy-tailed distribution in low-dim
- Stochastic optimization

charts/tsne\_umap\_comparison.pdf

### UMAP (Uniform Manifold Approximation)

- Faster than t-SNE
- Preserves global structure better

## Internal Metrics

Silhouette Score:

$$s(i) = \frac{b(i) - a(i)}{\max(a(i), b(i))}$$

where  $a(i)$  = avg distance within cluster,  $b(i)$  = avg distance to nearest cluster

Calinski-Harabasz Index:

$$CH = \frac{SS_B/(k-1)}{SS_W/(n-k)}$$

Davies-Bouldin Index:

$$DB = \frac{1}{k} \sum_{i=1}^k \max_{j \neq i} \frac{\sigma_i + \sigma_j}{d(c_i, c_j)}$$

Inertia (Within-cluster sum of squares):

$$WCSS = \sum_{i=1}^k \sum_{x \in C_i} \|x - \mu_i\|^2$$

## External Metrics

Adjusted Rand Index:

$$ARI = \frac{RI - E[RI]}{\max(RI) - E[RI]}$$

Normalized Mutual Information:

$$NMI = \frac{MI(U, V)}{\sqrt{H(U)H(V)}}$$

Homogeneity and Completeness:

- Homogeneity: Each cluster contains only one class
- Completeness: All members of class in same cluster

V-measure: Harmonic mean of homogeneity and completeness

**Note:** External metrics require ground truth labels

## Clustering

### Customer Segmentation:

- Group customers by behavior
- Targeted marketing campaigns
- Product recommendations

### Market Basket Analysis:

- Find product associations
- Store layout optimization
- Cross-selling opportunities

### Image Segmentation:

- Medical image analysis
- Computer vision
- Object recognition

## Dimensionality Reduction

### Data Visualization:

- Explore high-dimensional data
- Identify patterns and outliers
- Present insights to stakeholders

### Feature Engineering:

- Reduce computational cost
- Remove noise and redundancy
- Improve model performance

### Compression:

- Image and audio compression
- Efficient data storage
- Fast data transmission

## Anomaly Detection

### Fraud Detection:

- Credit card transactions
- Insurance claims
- Online account activity

### Network Security:

- Intrusion detection
- Malware identification
- Unusual traffic patterns

### Quality Control:

- Manufacturing defects
- System monitoring
- Predictive maintenance

Unsupervised learning reveals hidden patterns and structures in data without labeled examples

## Part 4: Neural Networks and Deep Learning

### From Perceptrons to Modern Architectures

## Mathematical Model

Linear Combination:

$$z = w_1x_1 + w_2x_2 + \dots + w_nx_n + b$$

$$z = \mathbf{w}^T \mathbf{x} + b$$

Activation Function:

$$y = \sigma(z) = \begin{cases} 1 & \text{if } z \geq 0 \\ 0 & \text{if } z < 0 \end{cases}$$

[charts/perceptron\\_model.pdf](#)

Decision Boundary:

$$\mathbf{w}^T \mathbf{x} + b = 0$$

Learning Rule (Perceptron Algorithm):

$$w_i := w_i + \eta(y - \hat{y})x_i$$

$$b := b + \eta(y - \hat{y})$$

where  $\eta$  is the learning rate

### Perceptron Limitations:

- Can only learn linearly separable functions
- Cannot solve XOR problem

## Architecture

Forward Propagation:

$$z^{[l]} = W^{[l]} a^{[l-1]} + b^{[l]}$$

$$a^{[l]} = g^{[l]}(z^{[l]})$$

**Universal Approximation Theorem:** A neural network with:

- One hidden layer
- Finite number of neurons
- Non-linear activation function

can approximate any continuous function on a compact set to arbitrary accuracy.

**Key Insight:** Width vs depth tradeoff

- Wide shallow networks: Exponential width needed
- Deep narrow networks: Polynomial parameters

[charts/mlp\\_architecture.pdf](#)

### Activation Functions:

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



## Algorithm

### Chain Rule Application:

$$\frac{\partial L}{\partial W^{[l]}} = \frac{\partial L}{\partial z^{[l]}} \frac{\partial z^{[l]}}{\partial W^{[l]}}$$

### Backward Pass:

$$\delta^{[l]} = \frac{\partial L}{\partial z^{[l]}}$$

$$\delta^{[l-1]} = (W^{[l]})^T \delta^{[l]} \odot g'(z^{[l-1]})$$

### Parameter Updates:

$$\frac{\partial L}{\partial W^{[l]}} = \delta^{[l]} (a^{[l-1]})^T$$

$$\frac{\partial L}{\partial b^{[l]}} = \delta^{[l]}$$

### Gradient Descent:

charts/backpropagation.pdf

### Computational Graph:

- Forward pass: Compute outputs
- Backward pass: Compute gradients

## Common Activations

Sigmoid:

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

Range:  $(0, 1)$ , smooth, vanishing gradients

Tanh:

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

Range:  $(-1, 1)$ , zero-centered, still vanishing gradients

ReLU:

$$\text{ReLU}(x) = \max(0, x)$$

Simple, fast, sparse activations, dying ReLU problem

Leaky ReLU:

$$\text{LeakyReLU}(x) = \max(\alpha x, x)$$

Fixes dying ReLU,  $\alpha = 0.01$  typically

`charts/activation_functions.pdf`

Modern Activations:

- **ELU:**  $f(x) = \begin{cases} x & x > 0 \\ \alpha(e^x - 1) & x \leq 0 \end{cases}$

## Architecture Components

### Convolution Operation:

$$(I * K)_{ij} = \sum_m \sum_n I_{i+m, j+n} K_{m,n}$$

### Key Concepts:

- **Local Connectivity:** Neurons connect to local regions
- **Parameter Sharing:** Same filter across all positions
- **Translation Invariance:** Features detected anywhere

### Typical CNN Architecture:

- ① Convolution + ReLU
- ② Pooling (max or average)
- ③ Repeat multiple times
- ④ Flatten and fully connected layers
- ⑤ Final classification layer

### Filter Parameters:

- Kernel size (3x3, 5x5, 7x7)
- Stride (step size)

charts/cnn\_architecture.pdf

### Pooling Operations:

- **Max Pooling:** Take maximum in region
- **Average Pooling:** Take average in region