ML Methods

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Introduction
  - Machine Learning
  - Motivation

A Practical View
  - Method or Models
  - Interpretability
  - Metric Choice

A Better Point of View
  - The Example of Univariate Linear Regression
  - Supervised Learning

Risk Estimation and Method Choice
  - Cross Validation
  - Cross Validation and Test
  - Cross Validation and Weights
  - Auto ML

A Probabilistic Point of View
  - Parametric Conditional Density Modeling
  - Non Parametric Conditional Density Modeling
  - Generative Modeling

Optimization Point of View
  - SVM
  - Penalization
  - (Deep) Neural Networks
  - Tree Based Methods
  - Ensemble Methods

Empirical Risk Minimization
  - Empirical Risk Minimization
  - ERM and PAC Bayesian Analysis
  - Hoeffding and Finite Class
  - McDiarmid and Rademacher Complexity
  - VC Dimension
  - Structural Risk Minimization

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Introduction

Machine Learning

Sources: MyCarDoesWhat.org/theverge.com/Rutgers Prep School
A definition by Tom Mitchell (http://www.cs.cmu.edu/~tom/)

A computer program is said to learn from experience $E$ with respect to some class of tasks $T$ and performance measure $P$, if its performance at tasks in $T$, as measured by $P$, improves with experience $E$. 
Object Detection

A detection algorithm:

- **Task**: say if an object is present or not in the image
- **Performance**: number of errors
- **Experience**: set of previously seen labeled images
An article clustering algorithm:

- **Task**: group articles corresponding to the same news
- **Performance**: quality of the clusters
- **Experience**: set of articles
A Robot that Learns

A robot endowed with a set of sensors playing football:

- **Task**: play football
- **Performance**: score evolution
- **Experience**:
  - past games
  - current environment and action outcome,
Three Kinds of Learning

Unsupervised Learning
- **Task:** Clustering/DR
- **Performance:** Quality
- **Experience:** Raw dataset (No Ground Truth)

Supervised Learning
- **Task:** Prediction/Classification
- **Performance:** Average error
- **Experience:** Good Predictions (Ground Truth)

Reinforcement Learning
- **Task:** Action
- **Performance:** Total reward
- **Experience:** Reward from env. (Interact. with env.)

**Timing:** Offline/Batch (learning from past data) vs Online (continuous learning)
Supervised and Unsupervised

**Supervised Learning (Imitation)**

**Goal:** Learn a function $f$ predicting a variable $Y$ from an individual $X$.

**Data:** Learning set with labeled examples $(X_i, Y_i)$

**Assumption:** Future data behaves as past data!

**Predicting is not explaining!**
Supervised and Unsupervised Learning

**Supervised Learning (Imitation)**
- **Goal:** Learn a function $f$ predicting a variable $Y$ from an individual $X$.
- **Data:** Learning set with labeled examples $(X_i, Y_i)$

**Assumption:** Future data behaves as past data!

**Predicting is not explaining!**

**Unsupervised Learning (Structure Discovery)**
- **Goal:** Discover a structure within a set of individuals $(X_i)$.
- **Data:** Learning set with unlabeled examples $(X_i)$

Unsupervised learning is not a well-posed setting...
Machine Can and Cannot

**Machine Can**
- Forecast (Prediction using the past)
- Detect some changes
- Memorize/Reproduce
- Take a decision very quickly
- Learn from huge dataset
- Optimize a single task
- Replace/Help some humans

**Machine Cannot**
- Predict something never seen before
- Detect any new behaviour
- Create something brand new
- Understand the world
- Get smart really fast
- Go beyond their task
- Kill all humans

- Some progresses but still very far from the *singularity*...
Machine Learning Methods

- Huge catalog of methods,
- Need to define the performance,
- Numerous tricks: feature design, hyperparameter selection...
Under and Over Fitting

Finding the Right Complexity

- What is best?
  - A simple model that is stable but false? *(oversimplification)*
  - A very complex model that could be correct but is unstable? *(conspiracy theory)*
  - Neither of them: tradeoff that depends on the dataset.
Introduction

Machine Learning Pipeline

Learning pipeline

- Test and compare models.
- Deployment pipeline is different!
Data Science ≠ Machine Learning

Main DS difficulties

- Figuring out the problem,
- Formalizing it,
- Storing and accessing the data,
- Deploying the solution,
- Not (always) the Machine Learning part!
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Reading a ZIP code on an envelop

- **Task:** give a number from an image.
- **Data:** $X =$ image / $Y =$ corresponding number.
- **Performance measure:** error rate.
Predicting protein interaction

- **Task:** Predict (unknown) interactions between proteins.
- **Data:** $X =$ pair of proteins / $Y =$ existence or no of interaction.
- **Performance measure:** error rate.
- Numerous similar questions in bio(informatics): genomic,…
Face detection

- **Task:** Detect the position of faces in an image
- Different setting?
- Reformulation as a supervised learning problem.
- **Goal:** Detect the presence of faces at several positions and scales.
- **Data:** $X =$ sub image / $Y =$ presence or no of a face...
- **Performance measure:** error rate.
- Lots of detections in an image: post processing required...
- **Performance measure:** box precision.
Height estimation

- Simple (and classical) dataset.
- **Task:** predict the height from circumference.
- **Data:** $X = \text{circumference} / \text{circ}$
- $Y = \text{height}$.
- **Performance measure:** means squared error.
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What is a Method?

A Standard Machine Learning Pipeline

A Learning Method

- Formula/Algorithm allowing to make predictions
- Algorithm allowing to chose this formula/algorithm
- Data preprocessing (cleansing, coding...)
- Optimization criterion for the choice!

Source: Microsoft
Simple Approach: Similarity

- Imitate the answer to give by mixing answers to similar questions (k nearest neighbors)
- Require to search for those similar questions for each request
- Not always very efficient but fast to build (less to use...)
- Easy to understand and rather stable
Simple Formula: Linear Method

Imitate the answer to give (linear regression) or a transformation of the conditional probability of the category (logistic regression)

Numerous variations on the parameter optimization (penalization, SVM, ...)

Pretty efficient and fast to build

Easy to understand and rather stable
Simple Algorithm: Tree

- Construction of a decision tree
- Impossible to really optimize but good tree can be obtained
- Not always very efficient but very quick to build
- Very easy to understand but not really stable
Ensemble Methods

Strategy:
- **Bagging**: construction of variations in parallel and averaging (**random forest**)
- **Boosting**: construction of sequential improvements (**XGBoost, Lightgbm**)
- **Stacking**: Use of a first set of predictors as features

- Very good performance for structured data but quite slow to build
- Stable but hard to understand
Deep Learning

- Chain of simple formulae (**Neural Network**)
- Joint optimization
- Very good performance for unstructured data but slow to build
- Mildly stable and very hard to understand
# Methods: Pros and Cons

<table>
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<tr>
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<th>Performance</th>
<th>Training Speed</th>
<th>Inf. Speed</th>
<th>Stability</th>
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## Take Away Message

- No unanimously best solution
- Impossible to guess which method is going to be the best!
- A good practice is to always try a linear method as well as an ensemble one for structured data or deep one for unstructured data
Preprocessing

- Art of creating sophisticated representations of initial data
- Key for good performances
- Examples: individual transformation, variable combination, category (and text) coding...

**Important part of the learning method**
**Methods/Models in Machine Learning**

- **ML Methods**
  - Huge catalog of methods,
  - Need to define the performance,
  - Need to represent well the data
  - Need to choose the best method yielding a good model
Under and Over Fitting

Finding the Right Complexity

- **What is best?**
  - A simple model that is stable but false? *(oversimplification)*
  - A very complex model that could be correct but is unstable? *(conspiracy theory)*
- Neither of them: tradeoff that depends on the dataset.
Which Method to Use?

Competition between several polynomial models.

- Toy model where everything is known.
Over-fitting

Model Prediction Error vs Model Complexity

Training Error vs Optimization

Prediction Error for New Data

Source: A. Ng
Learning pipeline

- Test and compare models.
- Deployment pipeline is different!
Cross Validation Principle

- Train a model and check its quality on different pieces of the data.

- Check the quality of a method by repeating the previous approach.

- **Beware:** a different predictor is learnt for each split.

Source: M. Kühn
The Full Cross Validation Scheme

- Most important part of machine learning.
- Automatic choice of model possible by (intelligent ?) exploration...
Best Polynomial

Competition results

- The true model is not the winner!
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Is this that easy?

- Simple formula setting:
  \[ Y \approx f(X) = a_0 + a_1 X^{(1)} + a_2 X^{(2)} + \cdots + a_d X^{(d)} \]

- Beware of the interpretation!

- Everything being equal... Correlation is not causality...
Interpretability or Explainability

- Interpretability: possibility to give a causal aspect to the formula.
- Explainability: possibility to find the variables having an effect on the decision and their effect.

- Explainability is much easier than interpretability.
- Transparency (on the datasets, the criterion optimized and the algorithms) yields already a lot of information.
A few directions

- Data Explanation.
- Use of explainable methods (linear?).
- Use of black box methods:
  - Global explanation (variable importance)
  - Local explanation (linear approximation, alternative scenario...)

- Causality very hard to access without a real experimental plan with interventions!
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Quality metric has a strong impact on the solution.

- Implicite encoding rather than an explicit one!
- Often simplified criterion in the optimization part.
- More involved criterion can be used in evaluation.
Supervised Performance Metrics

Measure of the cost of not being perfect!

- Criterion used to *optimize* the predictor and/or *evaluate* its interest.
- Classical metrics: quadratic error, zero/one error.
- Many other possible choices, ideally encoding domain expertise (asymmetry...)
- The criterion can be different between optimization and evaluation because of computation requirements.
- Very important factor (too) often neglected.
Unsupervised Performance Metrics

Measure the quality of the result!

- Dimension Reduction / Representation: reconstruction quality, relationship preservation.
- Clustering: measure of intra-group proximity and inter-group difference?
- Very subjective criterion!
- Hard to define the right distances especially for discrete variables.
- In practice, quality often evaluated by the a posteriori interest.
Fairness

Very hard to specify criterion.

No consensus on its definition:
  - faithful reproduction of the reality?
  - correction of its bias?

Current approaches through constraints in the optimization.

A posteriori verification unavoidable!
What About the Data Bias?

Central assumption: representativity of the data!

- Optimization made in this setting.
- Possible training data bias:
  - selection bias in the data
  - population evolution
  - (historical) bias in the targets
- Correction possible at least up to a certain point for the 2 first cases if one is aware of the situation.
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Simple (and classical) dataset.

Goal: predict the height from circumference

\[ X = \text{circ} = \text{circumference}. \]

\[ Y = \text{ht} = \text{height}. \]
**Linear Model**

- **Parametric model:**
  \[ f_\beta(circ) = \beta^{(1)} + \beta^{(2)} \times circ \]

- How to choose \( \beta = (\beta^{(1)}, \beta^{(2)}) \)?
Least Squares

Methodology

- Natural goodness criterion:
  \[
  \sum_{i=1}^{n} |Y_i - f_\beta(X_i)|^2 = \sum_{i=1}^{n} |ht_i - f_\beta(circ_i)|^2 \\
  = \sum_{i=1}^{n} |ht_i - (\beta^{(1)} + \beta^{(2)} circ_i)|^2
  \]

- Choice of $\beta$ that minimizes this criterion!
  \[
  \hat{\beta} = \arg\min_{\beta \in \mathbb{R}^2} \sum_{i=1}^{n} |h_i - (\beta^{(1)} + \beta^{(2)} circ_i)|^2
  \]

- Easy minimization with an explicit solution!
Prediction

- Linear prediction for the height:

\[ \hat{ht} = f_\beta(circ) = \hat{\beta}^{(1)} + \hat{\beta}^{(2)} \cdot circ \]
Heuristic

Linear Regression

- **Statistical model**: \((\text{circ}_i, \text{ht}_i)\) i.i.d. with the same law as a generic \((\text{circ}, \text{ht})\).
- **Performance criterion**: Look for \(f\) with a small average error
  \[ \mathbb{E}[|\text{ht} - f(\text{circ})|^2] \]
- **Empirical criterion**: Replace the unknown law by its empirical counterpart
  \[ \frac{1}{n} \sum_{i=1}^{n} |\text{ht}_i - f(\text{circ}_i)|^2 \]
- **Predictor model**: As the minimum over all function is 0 (if all the \(\text{circ}_i\) are different), restrict to the linear functions \(f(\text{circ}) = \beta^{(1)} + \beta^{(2)}\text{circ}\) to avoid over-fitting.
- **Model fitting**: Explicit formula here.

- This model can be too simple!
Polynomial Regression

Polynomial Model

- Polynomial model: \( f_\beta(\text{circ}) = \sum_{l=1}^{p} \beta^{(l)} \text{circ}^{l-1} \)
  - Linear in \( \beta \)!
  - Easy least squares estimation for any degree!
Which Degree?

Models

- Increasing degree = increasing complexity and better fit on the data
Which Degree?

Models

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Which Degree?

**Models**

- Increasing degree = increasing complexity and better fit on the data
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Models

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Which Degree?

## Models

- Increasing degree = increasing complexity and better fit on the data
Which Degree?

Best Degree?

- How to choose among those solutions?
Risk behavior

- Training error (empirical error on the training set) decays when the complexity of the model increases.
- Quite different behavior when the error is computed on new observations (true risk / generalization error).
- Overfit for complex models: parameters learned are too specific to the learning set!
- General situation! (Think of polynomial fit...)
- Need to use another criterion than the training error!
Cross Validation and Penalization

Two directions

- **How to estimate** the generalization error differently?
- **Find** a way to **correct** the empirical error?

Two Approaches

- **Cross validation:** Estimate the error on a different dataset:
  - Very efficient (and almost always used in practice!)
  - Need more data for the error computation.
- **Penalization approach:** Correct the optimism of the empirical error:
  - Require to find the correction (penalty).
Questions

- How to build a model?
- How to fit a model to the data?
- How to assess its quality?
- How to select a model among a collection?
- How to guaranty the quality of the selected model?
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Supervised Learning

Supervised Learning Framework

- Input measurement \( X \in \mathcal{X} \)
- Output measurement \( Y \in \mathcal{Y} \).
- \((X, Y) \sim \mathbb{P}\) with \(\mathbb{P}\) unknown.
- **Training data** : \( D_n = \{(X_1, Y_1), \ldots, (X_n, Y_n)\} \) (i.i.d. \(\sim \mathbb{P}\))

- Often
  - \( X \in \mathbb{R}^d \) and \( Y \in \{-1, 1\} \) (classification)
  - or \( X \in \mathbb{R}^d \) and \( Y \in \mathbb{R} \) (regression).
- A **predictor** is a function in \( \mathcal{F} = \{ f : \mathcal{X} \to \mathcal{Y} \text{ meas.} \} \)

Goal

- Construct a **good** predictor \( \hat{f} \) from the training data.

- Need to specify the meaning of good.
- Classification and regression are almost the **same** problem!
## Loss and Probabilistic Framework

### Loss function for a generic predictor

- **Loss function**: \( \ell(Y, f(X)) \) measures the goodness of the prediction of \( Y \) by \( f(X) \)
- **Examples**:
  - Prediction loss: \( \ell(Y, f(X)) = 1_{Y \neq f(X)} \)
  - Quadratic loss: \( \ell(Y, f(X)) = |Y - f(X)|^2 \)

### Risk function

- Risk measured as the average loss for a new couple:
  \[
  \mathcal{R}(f) = \mathbb{E}_{(X, Y) \sim P}[\ell(Y, f(X))]
  \]
- **Examples**:
  - Prediction loss: \( \mathbb{E}[\ell(Y, f(X))] = \mathbb{P}(Y \neq f(X)) \)
  - Quadratic loss: \( \mathbb{E}[\ell(Y, f(X))] = \mathbb{E}[|Y - f(X)|^2] \)
- **Beware**: As \( \hat{f} \) depends on \( D_n \), \( \mathcal{R}(\hat{f}) \) is a random variable!
Best Solution

- The best solution $f^*$ (which is independent of $D_n$) is
  \[ f^* = \arg\min_{f \in F} \mathcal{R}(f) = \arg\min_{f \in F} \mathbb{E}[\ell(Y, f(X))] = \arg\min_{f \in F} \mathbb{E}_X \left[ \mathbb{E}_{Y|X} [\ell(Y, f(X))] \right] \]

Bayes Predictor (explicit solution)

- In binary classification with $0-1$ loss:
  \[ f^*(X) = \begin{cases} +1 & \text{if } \mathbb{P}(Y = +1|X) \geq \mathbb{P}(Y = -1|X) \\ -1 & \text{otherwise} \end{cases} \]

- In regression with the quadratic loss
  \[ f^*(X) = \mathbb{E}[Y|X] \]

Issue: Solution requires to know $\mathbb{E}[Y|X]$ for all values of $X$!
Goal

Machine Learning

- Learn a rule to construct a predictor $\hat{f} \in \mathcal{F}$ from the training data $\mathcal{D}_n$ s.t. the risk $\mathcal{R}(\hat{f})$ is small on average or with high probability with respect to $\mathcal{D}_n$.

- In practice, the rule should be an algorithm!

Canonical example: Empirical Risk Minimizer

- One restricts $f$ to a subset of functions $S = \{f_\theta, \theta \in \Theta\}$
- One replaces the minimization of the average loss by the minimization of the empirical loss

$$\hat{f} = f_\hat{\theta} = \arg\min_{f_\theta, \theta \in \Theta} \frac{1}{n} \sum_{i=1}^{n} \ell(Y_i, f_\theta(X_i))$$

- Examples:
  - Linear regression
  - Linear classification with

$$S = \{ x \mapsto \text{sign} \{ x^T \beta + \beta^{(0)} \} / \beta \in \mathbb{R}^d, \beta^{(0)} \in \mathbb{R} \}$$
Example: TwoClass Dataset

Synthetic Dataset

- Two features/covariates.
- Two classes.

- Dataset from *Applied Predictive Modeling*, M. Kuhn and K. Johnson, Springer
- Numerical experiments with R and the `{caret}` package.
Example: Linear Discrimination

Logistic

Decision region

Decision boundary
Example: More Complex Model

Naive Bayes with kernel density estimates

Decision region

Decision boundary

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<th>PredictorB</th>
<th>PredictorA</th>
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<tr>
<td>0.2</td>
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classes
- Class1
- Class2

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classes
- Class1
- Class2
Dataset - P.A. Cornillon

- Real dataset of 1429 eucalyptus obtained by P.A. Cornillon:
  - $X$: circumference / $Y$: height
Dataset - P.A. Cornillon

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- Real dataset of 1429 eucalyptus obtained by P.A. Cornillon:
  - $X$: circumference, block, clone / $Y$: height
Under-fitting / Over-fitting Issue

Model Complexity Dilemma

- What is best a simple or a complex model?
- Too simple to be good? Too complex to be learned?
Under-fitting / Over-fitting Issue

- **Under-fitting**: simple model are too simple.
- **Over-fitting**: complex model are too specific to the training set.
Bias-Variance Dilemma

- General setting:
  - $\mathcal{F} = \{\text{measurable functions } \mathcal{X} \rightarrow \mathcal{Y}\}$
  - Best solution: $f^* = \arg\min_{f \in \mathcal{F}} \mathcal{R}(f)$
  - Class $S \subset \mathcal{F}$ of functions
  - Ideal target in $S$: $f^*_S = \arg\min_{f \in S} \mathcal{R}(f)$
  - Estimate in $S$: $\hat{f}_S$ obtained with some procedure

Approximation error and estimation error (Bias/ Variance)

$$\mathcal{R}(\hat{f}_S) - \mathcal{R}(f^*) = \underbrace{\mathcal{R}(f^*_S) - \mathcal{R}(f^*)}_{\text{Approximation error}} + \underbrace{\mathcal{R}(\hat{f}_S) - \mathcal{R}(f^*_S)}_{\text{Estimation error}}$$

- Approx. error can be large if the model $S$ is not suitable.
- Estimation error can be large if the model is complex.

Agnostic approach

- No assumption (so far) on the law of $(X, Y)$. 

A Better Point of View
Different behavior for different model complexity

- **Low complexity model** are easily learned but the approximation error *(bias)* may be large *(Under-fit)*.

- **High complexity model** may contain a good ideal target but the estimation error *(variance)* can be large *(Over-fit)*

**Bias-variance trade-off** ⇔ avoid overfitting and underfitting

- **Rk**: Better to think in term of method (including feature engineering and specific algorithm) rather than only of model.
Theoretical Analysis

Statistical Learning Analysis

- Error decomposition:
  \[
  R(\hat{f}_S) - R(f^*) = R(f^*_S) - R(f^*) + R(\hat{f}_S) - R(f^*_S)
  \]
  Approximation error  Estimation error

- Bound on the approximation term: approximation theory.
- Probabilistic bound on the estimation term: probability theory!
- **Goal: Agnostic bounds**, i.e. bounds that do not require assumptions on \( \mathbb{P} \)!
  (Statistical Learning?)

- Often need mild assumptions on \( \mathbb{P} \)… (Nonparametric Statistics?)
Binary Classification Loss Issue

Empirical Risk Minimizer

\[ \hat{f} = \arg\min_{f \in \mathcal{S}} \frac{1}{n} \sum_{i=1}^{n} \ell^{0/1}(Y_i, f(X_i)) \]

- Classification loss: \( \ell^{0/1}(y, f(x)) = 1_{y \neq f(x)} \)
- Not convex and not smooth!
The best solution $f^*$ (which is independent of $D_n$) is

$$f^* = \arg \min_{f \in \mathcal{F}} \mathbb{E}[\ell(Y, f(X))] = \arg \min_{f \in \mathcal{F}} \mathbb{E}_X [\mathbb{E}_{Y|X}[\ell(Y, f(X))]]$$

### Bayes Predictor (explicit solution)

In binary classification with $0−1$ loss:

$$f^*(X) = \begin{cases} +1 & \text{if } \mathbb{P}(Y = +1|X) \geq \mathbb{P}(Y = -1|X) \\ -1 & \text{otherwise} \end{cases}$$

**Issue:** Solution requires to know $\mathbb{E}[Y|X]$ for all values of $X$!

**Solution:** Replace it by an estimate.
Minimizer of the risk

\[ \hat{f} = \arg\min_{f \in S} \frac{1}{n} \sum_{i=1}^{n} \ell^{0/1}(Y_i, f(X_i)) \]

- **Issue:** Classification loss is not convex or smooth.
- **Solution:** Replace it by a convex majorant.
Probabilistic and Optimization Framework

How to find a good function $f$ with a small risk

$$ R(f) = \mathbb{E}[\ell(Y, f(X))] \quad ? $$

Canonical approach: $\hat{f}_S = \arg\min_{f \in S} \frac{1}{n} \sum_{i=1}^{n} \ell(Y_i, f(X_i))$

Problems

- How to choose $S$?
- How to compute the minimization?

A Probabilistic Point of View

Solution: For $X$, estimate $Y|X$ plug this estimate in the Bayes classifier:
(Generalized) Linear Models, Kernel methods, $k$-nn, Naive Bayes, Tree, Bagging...

An Optimization Point of View

Solution: If necessary replace the loss $\ell$ by an upper bound $\bar{\ell}$ and minimize the empirical loss: SVR, SVM, Neural Network, Tree, Boosting...
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   - Cross Validation and Weights
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   - Non Parametric Conditional Density Modeling
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   - Ensemble Methods

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   - McDiarmid and Rademacher Complexity
   - VC Dimension
   - Structural Risk Minimization

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![Data Scatter Plot](image)
Example: Linear Discrimination

Logistic

Decision region

Decision boundary

classes

Class1
Class2
Example: More Complex Model

Naive Bayes with kernel density estimates

Decision region

Decision boundary

classes

- Class1
- Class2
Example: KNN

k-NN with \( k=1 \)

Decision region

Decision boundary

- **classes**
  - Red: Class1
  - Blue: Class2
Example: KNN

k-NN with $k=5$

Decision region

Decision boundary
Example: KNN

k-NN with k=9

Decision region

Decision boundary

classes
- Class1
- Class2
Example: KNN

k-NN with k=13

Decision region

Decision boundary

classes

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Class1

Class2

classes

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Class1

Class2
Example: KNN

k-NN with $k=17$

Decision region

Decision boundary

classes

- Class1
- Class2

Risk Estimation and Method Choice
Example: KNN
Example: KNN

k-NN with k=25

Decision region

classes

Class1
Class2

Decision boundary

classes

Class1
Class2
Example: KNN

k-NN with $k=29$

Decision region

Decision boundary

classes

Class1

Class2
Example: KNN

k-NN with $k=33$

Decision region

Decision boundary

classes
- Class1
- Class2
Example: KNN

k-NN with k=37

Decision region

Decision boundary

classes
- Class1
- Class2
Example: KNN

k-NN with \( k=45 \)

Decision region

Decision boundary

classes
- Class1
- Class2

classes
- Class1
- Class2
Example: KNN

k-NN with k=53

Decision region

Decision boundary

classes
Class1
Class2

classes
Class1
Class2
Example: KNN

k-NN with $k=61$

Decision region:

- Classes: Class1 (red), Class2 (blue)

Decision boundary:

- Classes: Class1 (red), Class2 (blue)
Example: KNN

$k$-NN with $k=69$

Decision region

Decision boundary

classes

Class1

Class2

classes

Class1

Class2
Example: KNN

k-NN with k=77

Decision region

Decision boundary

classes

- Class1
- Class2
Example: KNN
Example: KNN

k-NN with $k=101$

Decision region

Decision boundary

classes
Class 1
Class 2

classes
Class 1
Class 2
Example: KNN

k-NN with $k=109$

Decision region

Decision boundary

- Classes: Class1, Class2

PredictorA

PredictorB
Example: KNN

k-NN with $k=117$

Decision region

Decision boundary
Example: KNN

k-NN with k=125

Decision region

Decision boundary

classes

Class1

Class2

classes

Class1

Class2
Example: KNN

k-NN with k=133

Decision region

Decision boundary

classes
Class1
Class2

classes
Class1
Class2
Example: KNN

k-NN with k=141

Decision region

Decision boundary
Example: KNN

$k$-NN with $k=149$

Decision region

Decision boundary

Classes:
- Class 1: Red
- Class 2: Blue
Example: KNN

k-NN with $k=157$

**Decision region**

- **PredictorA**
  - 0.2 to 0.6

- **PredictorB**
  - 0.2 to 0.6

**Classes**
- **Class 1** (Red)
- **Class 2** (Blue)

**Decision boundary**

- **PredictorA**
  - 0.2 to 0.6

- **PredictorB**
  - 0.2 to 0.6

**Classes**
- **Class 1** (Red)
- **Class 2** (Blue)
Example: KNN

k-NN with $k=165$

Decision region

Decision boundary
Example: KNN

k-NN with k=173

Decision region

Decision boundary

classes
- Class1
- Class2

classes
- Class1
- Class2
Example: KNN

k-NN with $k=181$

Decision region

Decision boundary

classes
- Class1
- Class2
Example: KNN

k-NN with k=189

Decision region

Decision boundary

classes

Class1

Class2
Example: KNN

k-NN with $k=197$

Decision region

Decision boundary
Learning/training risk (empirical risk on the learning/training set) decays when the complexity of the **method** increases.

Quite different behavior when the risk is computed on new observations (generalization risk).

Overfit for complex methods: parameters learned are too specific to the learning set!

General situation! (Think of polynomial fit...)  
Need to use a different criterion than the training risk!
Risk Estimation vs Method Selection

 Predictor Risk Estimation

- **Goal:** Given a predictor $f$ assess its quality.
- **Method:** Hold-out risk computation (Empirical risk correction).
- **Usage:** Compute an estimate of the risk of a selected $f$ using a test set to be used to monitor it in the future.

- Basic block very well understood.

 Method Selection

- **Goal:** Given a ML method assess its quality.
- **Method:** Cross Validation (Empirical risk correction)
- **Usage:** Compute risk estimates for several ML methods using training/validation sets to choose the most promising one.

- Estimates can be pointwise or better intervals.
- Multiple test issues in method selection.
Cross Validation and Empirical Risk Correction

Two Approaches

- **Cross validation:** Use empirical risk criterion but on independent data, very efficient (and almost always used in practice!) but slightly biased as its target uses only a fraction of the data.

- **Correction approach:** use empirical risk criterion but correct it with a term increasing with the complexity of $S$

\[ R_n(f_S) \rightarrow R_n(f_S) + \text{cor}(S) \]

and choose the method with the smallest corrected risk.

Which loss to use?

- The loss used in the risk: most natural!

- The loss used to estimate $\hat{\theta}$: penalized estimation!

- Other performance measure can be used.
• **Very simple idea:** use a second learning/verification set to compute a verification risk.
• Sufficient to remove the dependency issue!
• Implicit random design setting...

Cross Validation

- Use \((1 - \epsilon) \times n\) observations to train and \(\epsilon \times n\) to verify!
- Possible issues:
  - Validation for a learning set of size \((1 - \epsilon) \times n\) instead of \(n\) ?
  - Unstable risk estimate if \(\epsilon n\) is too small ?

- Most classical variations:
  - Hold Out,
  - Leave One Out,
  - \(V\)-fold cross validation.
Hold Out

**Principle**
- Split the dataset $\mathcal{D}$ in 2 sets $\mathcal{D}_{\text{train}}$ and $\mathcal{D}_{\text{test}}$ of size $n \times (1 - \epsilon)$ and $n \times \epsilon$.
- Learn $\hat{f}^{HO}$ from the subset $\mathcal{D}_{\text{train}}$.
- Compute the empirical risk on the subset $\mathcal{D}_{\text{test}}$:
  \[
  R_{n}^{HO}(\hat{f}^{HO}) = \frac{1}{n\epsilon} \sum_{(X_i, Y_i) \in \mathcal{D}_{\text{test}}} \ell(Y_i, \hat{f}^{HO}(X_i))
  \]

**Predictor Risk Estimation**
- Use $\hat{f}^{HO}$ as predictor.
- Use $R_{n}^{HO}(\hat{f}^{HO})$ as an estimate of the risk of this estimator.

**Method Selection by Cross Validation**
- Compute $R_{n}^{HO}(\hat{f}_{S}^{HO})$ for all the considered methods,
- Select the method with the smallest CV risk,
- Reestimate the $\hat{f}_{S}$ with all the data.
Hold Out

**Principle**

- Split the dataset $\mathcal{D}$ in 2 sets $\mathcal{D}_{\text{train}}$ and $\mathcal{D}_{\text{test}}$ of size $n \times (1 - \epsilon)$ and $n \times \epsilon$.
- Learn $\hat{f}^{\text{HO}}$ from the subset $\mathcal{D}_{\text{train}}$.
- Compute the empirical risk on the subset $\mathcal{D}_{\text{test}}$:
  \[
  R_n^{\text{HO}}(\hat{f}^{\text{HO}}) = \frac{1}{n\epsilon} \sum_{(X_i, Y_i) \in \mathcal{D}_{\text{test}}} \ell(Y_i, \hat{f}^{\text{HO}}(X_i))
  \]

- Only possible setting for risk estimation.

**Hold Out Limitation for Method Selection**

- Biased toward simpler method as the estimation does not use all the data initially.
- Learning variability of $R_n^{\text{HO}}(\hat{f}^{\text{HO}})$ not taken into account.
**V-fold Cross Validation**

### Principle

- **Split the dataset** $\mathcal{D}$ **in** $V$ **sets** $\mathcal{D}_v$ **of almost equals size.**
- **For** $v \in \{1, \ldots, V\}$:
  - **Learn** $\hat{f}^{-v}$ **from the dataset** $\mathcal{D}$ **minus the set** $\mathcal{D}_v$.
  - **Compute the empirical risk:**
    \[
    R_n^{-v}(\hat{f}^{-v}) = \frac{1}{n_v} \sum_{(X_i, Y_i) \in \mathcal{D}_v} \ell(Y_i, \hat{f}^{-v}(X_i))
    \]
  - **Compute the average empirical risk:**
    \[
    R_n^{CV}(\hat{f}) = \frac{1}{V} \sum_{v=1}^{V} R_n^{-v}(\hat{f}^{-v})
    \]
- **Estimation of the quality of a method not of a given predictor.**
- **Leave One Out** : $V = n$. 

---

Source: M. Kühn
\( V \)-fold Cross Validation

**Analysis (when \( n \) is a multiple of \( V \))**

- The \( R_n^{-V}(\hat{f}^{-V}) \) are identically distributed variable but are not independent!
- Consequence:

\[
\begin{align*}
\mathbb{E} \left[ R_n^{CV}(\hat{f}) \right] &= \mathbb{E} \left[ R_n^{-V}(\hat{f}^{-V}) \right] \\
\text{Var} \left[ R_n^{CV}(\hat{f}) \right] &= \frac{1}{V} \text{Var} \left[ R_n^{-V}(\hat{f}^{-V}) \right] \\
&\quad + \left( 1 - \frac{1}{V} \right) \text{Cov} \left[ R_n^{-V}(\hat{f}^{-V}), R_n^{-V}(\hat{f}^{-V}') \right]
\end{align*}
\]

- Average risk for a sample of size \( (1 - \frac{1}{V})n \).
- Variance term much more complex to analyze!
- Fine analysis shows that the larger \( V \) the better...

- Accuracy/Speed tradeoff: \( V = 5 \) or \( V = 10 \)!
Linear Regression and Leave One Out

- Leave One Out = V fold for $V = n$: very expensive in general.

A fast LOO formula for the linear regression

- **Prop:** for the least squares linear regression,
  \[
  \hat{f}^{-i}(X_i) = \frac{\hat{f}(X_i) - h_{ii}Y_i}{1 - h_{ii}}
  \]
  with $h_{ii}$ the $i$th diagonal coefficient of the hat (projection) matrix.

- Proof based on linear algebra!

- Leads to a fast formula for LOO:
  \[
  R_n^{LOO}(\hat{f}) = \frac{1}{n} \sum_{i=1}^{n} \frac{|Y_i - \hat{f}(X_i)|^2}{(1 - h_{ii})^2}
  \]
Cross Validation

Risk Estimation and Method Choice
Example: KNN ($\hat{k} = 61$ using cross-validation)
Selection Bias Issue:

- After method selection, the cross validation is biased.
- Furthermore, it qualifies the method and not the final predictor.
- Need to (re)estimate the risk of the final predictor.

(Train/Validation)/Test strategy

- **Split** the dataset in two a (Train/Validation) and Test.
- Use **CV** with the (Train/Validation) to **select a method**.
- Train this method on (Train/Validation) to **obtain a single predictor**.
- Estimate the **performance of this predictor** on Test.

Every choice made from the data is part of the method!
Risk Correction

- Empirical loss of an estimator computed on the dataset used to chose it is biased!
- Empirical loss is an optimistic estimate of the true loss.

**Risk Correction Heuristic**

- Estimate an upper bound of this optimism for a given family.
- Correct the empirical loss by adding this upper bound.

**Rk:** Finding such an upper bound can be complicated!
- Correction often called a **penalty**.
Penalization

Penalized Loss

- Minimization of
  \[
  \argmin_{\theta \in \Theta} \frac{1}{n} \sum_{i=1}^{n} \ell(Y_i, f_{\theta}(X_i)) + \text{pen}(\theta)
  \]

where \(\text{pen}(\theta)\) is a risk correction (penalty).

Penalties

- Upper bound of the optimism of the empirical loss
- Depends on the loss and the framework!

Instantiation

- Mallows Cp: Least Squares with \(\text{pen}(\theta) = 2\frac{d}{n}\sigma^2\).
- AIC Heuristics: Maximum Likelihood with \(\text{pen}(\theta) = \frac{d}{n}\).
- BIC Heuristics: Maximum Likelihood with \(\text{pen}(\theta) = \log(n)\frac{d}{n}\).
- Structural Risk Minimization: Pred. loss and clever penalty.
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Comparison of Two Means

Means

- **Setting:** r.v. $e_i^{(l)}$ with $1 \leq i \leq n_l$ and $l \in \{1, 2\}$ and their means
  \[
  \overline{e^{(l)}} = \frac{1}{n_l} \sum_{i=1}^{n_l} e_i^{(l)}
  \]

- **Question:** are the means $\overline{e^{(l)}}$ statistically different?

Classical i.i.d setting

- **Assumption:** $e_i^{(l)}$ are i.i.d. for each $l$.
- **Test formulation:** Can we reject the null hypothesis that $E[e^{(1)}] = E[e^{(2)}]$?
- **Methods:**
  - Gaussian (Student) test using asymptotic normality of a mean.
  - Non-parametric permutation test.

- Gaussian approach is linked to confidence intervals.
- The larger $n_l$ the smaller the confidence intervals.
Comparison of Two Means

Non i.i.d. case

- **Assumption:** \( e_i^{(l)} \) are i.d. for each \( l \) but not necessarily independent.
- **Test formulation:** Can we reject the null hypothesis that \( \mathbb{E}[e^{(1)}] = \mathbb{E}[e^{(2)}] \)?
- **Methods:**
  - Gaussian (Student) test using asymptotic normality of a mean but variance is hard to estimate.
  - Non-parametric permutation test but no confidence intervals.

- Setting for Cross Validation (other than holdout).
- Much more complicated than the i.i.d. case
Comparison of Several Means

Several means

- **Assumption:** $e_i^{(l)}$ are i.d. for each $l$ but not necessarily independent.

- **Tests formulation:**
  - Can we reject the null hypothesis that the $\mathbb{E}[e^{(l)}]$ are different?
  - Is the smaller mean statistically smaller than the second one?

- **Methods:**
  - Gaussian (Student) test using asymptotic normality of a mean with multiple tests correction.
  - Non-parametric permutation test but no confidence intervals.

- Setting for Cross Validation (other than holdout).
  - The more models one compares:
    - the larger the confidence intervals
    - the most probable the best model is a lucky winner
  - Justify the fallback to the simplest model that could be the best one.
PAC Approach

CV Risk, Methods and Predictors
- Cross-Validation risk: estimate of the average risk of a ML method.
- No risk bound on the predictor obtained in practice.

Probabilly-Approximately-Correct (PAC) Approach
- Replace the control on the average risk by a probabilistic bound
  \[ P\left(\mathbb{E}\left[\ell(Y, \hat{f}(X))\right] > R\right) \leq \epsilon \]
- Requires estimating quantiles of the risk.
Cross Validation and Confidence Interval

- How to replace pointwise estimation by a confidence interval?
- Can we use the variability of the CV estimates?
- **Negative result:** No unbiased estimate of the variance!

**Gaussian Interval (Comparison of the means and \sim indep.)**

- Compute the empirical variance and divide it by the number of folds to construct an asymptotic Gaussian confidence interval,
- Select the simplest model whose value falls into the confidence interval of the model having the smallest CV risk.

**PAC approach (Quantile, \sim indep. and small risk estim. error)**

- Compute the raw medians (or a larger raw quantiles)
- Select the model having the smallest quantiles to ensure a small risk with high probability.

- Always reestimate the chosen model with all the data.
- To obtain an unbiased risk estimate of the final predictor: hold out risk on untouched test data.
Cross Validation

variable
- AccuracyCV
- AccuracyCVInf
- AccuracyCVPAC
- Accuracy

model

value
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Unbalanced and Rebalanced Dataset

### Unbalanced Class
- **Setting:** One of the class is much more present than the other.
- **Issue:** Classifier *too attracted* by the majority class!

### Rebalanced Dataset
- **Setting:** Class proportions are different in the training and testing set (stratified sampling)
- **Issue:** Training risks are not estimate of testing risks.
Resampling Strategies

- Modify the training dataset so that the classes are more balanced.
- Two flavors:
  - Sub-sampling which spoils data,
  - Over-sampling which needs to create new examples.
- **Issues**: Training data is not anymore representative of testing data
- **Hard to do it right!**
## Estimation and Method Choice

### Testing
- **Testing class prob.:** \( \pi_t(k) \)
- **Testing risk target:**
  \[
  \mathbb{E}_{\pi_t}[\ell(Y, f(X))] = \sum_k \pi_t(k) \mathbb{E}[\ell(Y, f(X))| Y = k]
  \]

### Training
- **Training class prob.:** \( \pi_{tr}(k) \)
- **Training risk target:**
  \[
  \mathbb{E}_{\pi_{tr}}[\ell(Y, f(X))] = \sum_k \pi_{tr}(k) \mathbb{E}[\ell(Y, f(X))| Y = k]
  \]

### Implicit Testing Risk Using the Training One
- **Amounts to use a weighted loss:**
  \[
  \mathbb{E}_{\pi_{tr}}[\ell(Y, f(X))] = \sum_k \pi_{tr}(k) \mathbb{E}[\ell(Y, f(X))| Y = k]
  = \sum_k \pi_t(k) \mathbb{E} \left[ \frac{\pi_{tr}(k)}{\pi_t(k)} \ell(Y, f(X)) \middle| Y = k \right]
  = \mathbb{E}_{\pi_t} \left[ \frac{\pi_{tr}(Y)}{\pi_t(Y)} \ell(Y, f(X)) \right]
  \]
- **Put more weight on less probable classes!**
Weighted Loss

- In unbalanced situation, often the cost of misprediction is not the same for all classes (e.g. medical diagnosis, credit lending...)
- Much better to use this explicitly than to do blind resampling!

Weighted Loss

- Weighted loss:
  \[
  \ell(Y, f(X)) \rightarrow C(Y)\ell(Y, f(X))
  \]
- Weighted risk target:
  \[
  \mathbb{E}[C(Y)\ell(Y, f(X))]
  \]

- Rk: Strong link with \(\ell\) as \(C\) is independent of \(f\).
- Often allow reusing algorithm constructed for \(\ell\).
- \(C\) may also depend on \(X\)...
Weighted Loss, $\ell^{0/1}$ loss and Bayes Classifier

- The Bayes classifier is now:
  $$f^* = \arg\min \mathbb{E}[C(Y)\ell(Y, f(X))] = \arg\min \mathbb{E}_X[\mathbb{E}_{Y|X}[C(Y)\ell(Y, f(X))]]$$

**Bayes Predictor**

- For $\ell^{0/1}$ loss,
  $$f^*(X) = \arg\max_k C(k)\mathbb{P}(Y = k|X)$$

- Same effect than a threshold modification for the binary setting!
- Allow putting more emphasis on some classes than others.
Linking Weights and Proportions

Cost and Proportions

- Testing risk target:
  \[ \mathbb{E}_{\pi_t}[C_t(Y)\ell(Y, f(X))] = \sum_k \pi_t(k)C_t(k)\mathbb{E}[\ell(Y, f(X))|Y = k] \]

- Training risk target
  \[ \mathbb{E}_{\pi_{tr}}[C_{tr}(Y)\ell(Y, f(X))] = \sum_k \pi_{tr}(k)C_{tr}(k)\mathbb{E}[\ell(Y, f(X))|Y = k] \]

- **Coincide if**
  \[ \pi_t(k)C_t(k) = \pi_{tr}(k)C_{tr}(k) \]

- Lots of flexibility in the choice of \( C_t, \ C_{tr} \) or \( \pi_{tr} \)!
Combining Weights and Resampling

**Weighted Loss and Resampling**

- **Weighted loss:** choice of a weight $C_t \neq 1$.
- **Resampling:** use a $\pi_{tr} \neq \pi_t$.

Stratified sampling may be used to reduce the size of a dataset without losing a low probability class!

**Combining Weights and Resampling**

- **Weighted loss:** use $C_{tr} = C_t$ as $\pi_{tr} = \pi_t$.
- **Resampling:** use an implicit $C_t(k) = \pi_{tr}(k)/\pi_t(k)$.
- ** Combined:** use $C_{tr}(k) = C_t(k)\pi_t(k)/\pi_{tr}(k)$.

Most ML methods allow such weights!
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Auto ML

- Automatically propose a good predictor
- Rely heavily on risk evaluations
- **Pros:** easy way to obtain an excellent baseline
- **Cons:** black box that can be abused...
Auto ML Task

Input:
- a dataset $\mathcal{D} = (X_i, Y_i)$
- a loss function $\ell(Y, f(X))$
- a set of possible predictors $f_{l,h,\theta}$ corresponding to a method $l$ in a list, with hyperparameters $h$ and parameters $\theta$

Output:
- a predictor $f$ equal to $f_{\hat{l},\hat{h},\hat{\theta}}$ or combining several such functions.
Predictors, a.k.a fitted pipelines

- **Preprocessing:**
  - Feature design: normalization, coding, kernel...
  - Missing value strategy
  - Feature selection method

- **ML Method:**
  - Method itself
  - Hyperparameters and architecture
  - Fitted parameters (includes optimization algorithm)

- Quickly amounts to 20 to 50 design decisions!
- **Bruteforce exploration impossible!**
Auto ML and Hyperparameter Optimization

Most Classical Approach of Auto ML

- Task rephrased as an optimization on the discrete/continous space of methods/hyperparameters/parameters.
- Parameters obtained by classical minimization.
- Optimization of methods/hyperparameters much more challenging.
- Approaches:
  - Bruteforce: Grid search and random search
  - Clever exploration: Evolutionary algorithm
  - Surrogate based: Bayesian search and Reinforcement learning

Source: J. Borgstra
Learn from other Learning Tasks

- Consider the choice of the method from a dataset and a metric as a learning task.
- Requires a way to describe the problems (or to compute a similarity).
- Descriptor often based on a combination of dataset properties and fast method results.
- May output a list of candidates instead of a single method.

- Promising but still quite experimental!
How to obtain a good result with a time constraint?

- **Brute force**: Time out and methods screening with Meta-Learning (less exploration at the beginning)
- **Surrogate based**: Bayesian optimization (exploration/exploitation tradeoff)
- **Successive elimination**: Fast but not accurate performance evaluation at the beginning to eliminate the worst models (more exploration at the beginning)
- **Combined strategy**: Bandit strategy to obtain a more accurate estimate of risks only for the promising models (exploration/exploitation tradeoff)
Auto ML benchmark

Benchmark

- Not always (much) better than a good random forest or gradient boosting predictor.
- Worth the try!
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Three Classical Methods in a Nutshell

**Logistic Regression**
- Let \( f_\theta(X) = X^T \beta + \beta^{(0)} \) with \( \theta = (\beta, \beta^{(0)}) \).
- Let \( P_{\theta}(Y = 1|X) = e^{-f_\theta(X)}/(1 + e^{f_\theta(X)}) \)
- Estimate \( \theta \) by \( \hat{\theta} \) using a Maximum Likelihood.
- Classify using \( P_{\hat{\theta}}(Y = 1|X) > 1/2 \)

**\( k \) Nearest Neighbors**
- For any \( X' \), define \( V_{X'} \) as the \( k \) closest samples \( X_i \) from the dataset.
- Compute a score \( g_k = \sum_{X_i \in V_{X'}} 1_{Y_i = k} \)
- Classify using \( \text{arg max} g_k \) (majority vote).
Three Classical Methods in a Nutshell

**Quadratic Discrimant Analysis**

- For each class, estimate the mean $\mu_k$ and the covariance matrix $\Sigma_k$.
- Estimate the proportion $\mathbb{P}(Y = k)$ of each class.
- Compute a score $\ln(\mathbb{P}(X|Y = k)) + \ln(\mathbb{P}(Y = k))$

$$g_k(X) = -\frac{1}{2}(X - \mu_k)^\top \Sigma_k^{-1}(X - \mu_k)$$

$$-\frac{d}{2} \ln(2\pi) - \frac{1}{2} \ln(|\Sigma_k|) + \ln(\mathbb{P}(Y = k))$$

- Classify using $\text{arg max } g_k$

- Those three methods rely on a similar heuristic: the probabilistic point of view!
Best Solution

- The best solution \( f^* \) (which is independent of \( D_n \)) is
  \[
  f^* = \arg \min_{f \in \mathcal{F}} R(f) = \arg \min_{f \in \mathcal{F}} \mathbb{E}[\ell(Y, f(X))] = \arg \min_{f \in \mathcal{F}} \mathbb{E}_X \left[ \mathbb{E}_{Y|X}[\ell(Y, f(X))] \right]
  \]

Bayes Predictor (explicit solution)

- In binary classification with 0−1 loss:
  \[
  f^*(X) = \begin{cases} 
  +1 & \text{if } \mathbb{P}(Y = +1|X) \geq \mathbb{P}(Y = -1|X) \\
  \iff \mathbb{P}(Y = +1|X) \geq 1/2 \\
  -1 & \text{otherwise}
  \end{cases}
  \]

- In regression with the quadratic loss
  \[
  f^*(X) = \mathbb{E}[Y|X]
  \]

**Issue:** Explicit solution requires to know \( Y|X \) (or \( \mathbb{E}[Y|X] \)) for all values of \( X \)!
Plugin Predictor

- **Idea:** Estimate $Y|X$ by $\hat{Y}|X$ and plug it the Bayes classifier.

### Plugin Bayes Predictor

- In binary classification with $0 - 1$ loss:
  $$\hat{f}(X) = \begin{cases} +1 & \text{if } \frac{P(Y = +1|X)}{P(Y = -1|X)} \geq 1/2 \\ -1 & \text{otherwise} \end{cases}$$

- In regression with the quadratic loss
  $$\hat{f}(X) = E[Y|X]$$

- **Rk:** Direct estimation of $E[Y|X]$ by $\hat{E}[Y|X]$ also possible...
Plugin Predictor

- How to estimate $Y|X$?

**Three main heuristics**

- **Parametric Conditional modeling:** Estimate the law of $Y|X$ by a parametric law $L_\theta(X)$: (generalized) linear regression...

- **Non Parametric Conditional modeling:** Estimate the law of $Y|X$ by a non parametric estimate: kernel methods, loess, nearest neighbors...

- **Fully Generative modeling:** Estimate the law of $(X, Y)$ and use the Bayes formula to deduce an estimate of $Y|X$: LDA/QDA, Naive Bayes...

- **Rk:** Direct estimation of $\mathbb{E}[Y|X]$ by $\mathbb{E}[Y|X]$ also possible...
Plugin Classifier

- **Input**: a data set $D_n$
  
  Learn $Y|X$ or equivalently $P(Y = k|X)$ (using the data set) and plug this estimate in the Bayes classifier

- **Output**: a classifier $\hat{f} : \mathbb{R}^d \rightarrow \{-1, 1\}$

  
  $$\hat{f}(X) = \begin{cases} 
  +1 & \text{if } P(\hat{Y} = 1|X) \geq P(\hat{Y} = -1|X) \\
  -1 & \text{otherwise}
  \end{cases}$$

- Can we guarantee that the classifier is good if $Y|X$ is well estimated?
Classification Risk Analysis

**Theorem**

- If \( \hat{f} = \text{sign}(2\hat{p}_+ - 1) \) then
  \[
  \mathbb{E} \left[ \ell_{0,1}(Y, \hat{f}(X)) \right] - \mathbb{E} \left[ \ell_{0,1}(Y, f^*(X)) \right] \\
  \leq \mathbb{E} \left[ \| \hat{Y}|X - Y|X \|_1 \right] \\
  \leq \left( \mathbb{E} \left[ 2\text{KL}(Y|X, \hat{Y}|X) \right] \right)^{1/2}
  \]

- If one estimates \( \mathbb{P}(Y = 1|X) \) well then one estimates \( f^* \) well!
- Link between a *conditional density estimation* task and a *classification* one!
- **Rk:** In general, the conditional density estimation task is more complicated as one should be good for all values of \( \mathbb{P}(Y = 1|X) \) while the classification task focus on values around \( 1/2 \) for the \( 0/1 \) loss!
- In *regression*, (often) direct control of the quadratic loss...
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Parametric Conditional Density Models

- **Idea:** Estimate directly $Y|X$ by a parametric conditional density $P_\theta(Y|X)$.

**Maximum Likelihood Approach**

- Classical choice for $\theta$:
  \[
  \hat{\theta} = \arg\min_{\theta} - \sum_{i=1}^{n} \log P_\theta(Y_i|X_i)
  \]

- **Goal:** Minimize the Kullback-Leibler divergence between the conditional law of $Y|X$ and $P_\theta(Y|X)$
  \[
  E[KL(Y|X, P_\theta(Y|X))]
  \]

- **Rk:** This is often not (exactly) the learning task!

- Large choice for the family $\{P_\theta(Y|X)\}$ but depends on $Y$ (and $X$).

- **Regression:** One can also model directly $E[Y|X]$ by $f_\theta(X)$ and estimate it with a least square criterion...
Linear Conditional Density Models

Classical choice: \( \theta = (\theta', \varphi) \)

\[ P_\theta(Y|X) = P_{X^T \beta, \varphi}(Y) \]

Very strong assumption!

Classical examples:
- Binary variable: logistic, probit...
- Discrete variable: multinomial logistic regression...
- Integer variable: Poisson regression...
- Continuous variable: Gaussian regression...
Binary Classifier

Plugin Linear Classification

- Model $\mathbb{P}(Y = +1|X)$ by $h(X^T \beta + \beta^{(0)})$ with $h$ non decreasing.
- $h(X^T \beta + \beta^{(0)}) > 1/2 \iff X^T \beta + \beta^{(0)} - h^{-1}(1/2) > 0$
- Linear Classifier: $\text{sign}(X^T \beta + \beta^{(0)} - h^{-1}(1/2))$

Plugin Linear Classifier Estimation

- Classical choice for $h$:
  $$h(t) = \frac{e^t}{1 + e^t}$$ for logit or logistic
  $$h(t) = F_N(t)$$ for probit
  $$h(t) = 1 - e^{-e^t}$$ for log-log
- Choice of the best $\beta$ from the data.
Maximum Likelihood Estimate

Probabilistic Model

- By construction, $Y|X$ follows $B(\mathbb{P}(Y = +1|X))$
- Approximation of $Y|X$ by $B(h(x^\top \beta + \beta^{(0)}))$
- *Natural* probabilistic choice for $\beta$: maximum likelihood estimate.
- *Natural* probabilistic choice for $\beta$: $\beta$ approximately minimizing a distance between $B(h(x^\top \beta))$ and $B(\mathbb{P}(Y = 1|X))$.

Maximum Likelihood Approach

- Minimization of the negative log-likelihood:
  $$- \sum_{i=1}^{n} \log(\mathbb{P}(Y_i|X_i)) = - \sum_{i=1}^{n} \left(1_{Y_i=1} \log(h(X_i^\top \beta)) + 1_{Y_i=-1} \log(1 - h(X_i^\top \beta))\right)$$
- Minimization possible if $h$ is regular...
Maximum Likelihood Estimate

**KL Distance and negative log-likelihood**

- *Natural distance: Kullback-Leibler divergence*

\[
\text{KL}(\mathcal{B}(P(Y = 1|X)), \mathcal{B}(h(X^\top \beta))) = \mathbb{E}_X \left[ P(Y = 1|X) \log \frac{P(Y = 1|X)}{h(X^\top \beta)} \right.
\]

\[
\left. + P(Y = -1|X) \log \frac{1 - P(Y = 1|X)}{1 - h(X^\top \beta)} \right] = \mathbb{E}_X \left[ -P(Y = 1|X) \log(h(X^\top \beta)) \right.
\]

\[
\left. - P(Y = -1|X) \log(1 - h(X^\top \beta)) \right] + C_{X,Y}
\]

- *Empirical counterpart = negative log-likelihood (up to 1/n factor):*

\[
- \frac{1}{n} \sum_{i=1}^{n} \left( \mathbf{1}_{Y_i=1} \log(h(X_i^\top \beta)) + \mathbf{1}_{Y_i=-1} \log(1 - h(X_i^\top \beta)) \right)
\]
Logistic Regression

Logistic Regression and Odd

- Logistic model: \( h(t) = \frac{e^t}{1+e^t} \) (most natural choice...)
- The Bernoulli law \( B(h(t)) \) satisfies then
  \[
  \frac{P(Y = 1)}{P(Y = -1)} = e^t \iff \log \frac{P(Y = 1)}{P(Y = -1)} = t
  \]
- Interpretation in term of odd.
- Logistic model: linear model on the logarithm of the odd
  \[
  \log \frac{P(Y = 1|X)}{P(Y = -1|X)} = X^T \beta
  \]

Associated Classifier

- Plugin strategy:
  \[
  f_\beta(X) = \begin{cases} 
  1 & \text{if } \frac{e^{X^T \beta}}{1+e^{X^T \beta}} > 1/2 \iff X^T \beta > 0 \\
  -1 & \text{otherwise}
  \end{cases}
  \]
Likelihood Rewriting

- Negative log-likelihood:

\[- \frac{1}{n} \sum_{i=1}^{n} \left( 1_{Y_i=1} \log(h(X_i^T \beta)) + 1_{Y_i=-1} \log(1 - h(X_i^T \beta)) \right)\]

\[= - \frac{1}{n} \sum_{i=1}^{n} \left( 1_{Y_i=1} \log \left( \frac{e^{X_i^T \beta}}{1 + e^{X_i^T \beta}} \right) + 1_{Y_i=-1} \log \left( \frac{1}{1 + e^{X_i^T \beta}} \right) \right)\]

\[= \frac{1}{n} \sum_{i=1}^{n} \log \left( 1 + e^{-Y_i(X_i^T \beta)} \right)\]

- Convex and smooth function of $\beta$
- Easy optimization.
Example: Logistic
Feature Design

Transformed Representation
- From $X$ to $\Phi(X)$!
- New description of $X$ leads to a different \textbf{linear} model:
  $$f_\beta(X) = \Phi(X)^T \beta$$

Feature Design
- Art of choosing $\Phi$.
- Examples:
  - Renormalization, (domain specific) transform
  - Basis decomposition
  - Interaction between different variables...
Example: Quadratic Logistic
Gaussian Linear Regression

Gaussian Linear Model

- **Model:** \( Y|X \sim \mathcal{N}(X^\top \beta, \sigma^2) \) plus independence
- Probably the most classical model of all time!
- Maximum Likelihood with explicit formulas for the two parameters.

In regression, estimation of \( \mathbb{E}[Y|X] \) is sufficient: other/no model for the noise possible.
Extension of Gaussian Linear Regression

**Generalized Linear Model**

- Model entirely characterized by its mean (up to a scalar nuisance parameter) \( (v(E_\theta[Y]) = \theta \) with \( v \) invertible).
- Exponential family: Probability law family \( P_\theta \) such that the density can be written

\[
  f(y, \theta, \varphi) = e^{\frac{y\theta - v(\theta)}{\varphi} + w(y, \varphi)}
\]

where \( \varphi \) is a nuisance parameter and \( w \) a function independent of \( \theta \).

- Examples:
  - Gaussian: \( f(y, \theta, \varphi) = e^{-\frac{y\theta - \theta^2/2}{\varphi} - \frac{y^2/2}{\varphi}} \)
  - Bernoulli: \( f(y, \theta) = e^{y\theta - \ln(1 + e^{\theta})} \) (\( \theta = \ln \frac{p}{1 - p} \))
  - Poisson: \( f(y, \theta) = e^{(y\theta - e^{\theta}) + \ln(y!)} \) (\( \theta = \ln \lambda \))

- Linear Conditional model: \( Y|X \sim P_{x^T \beta} \ldots \)

- ML fit of the parameters
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Non Parametric Conditional Estimation

- **Idea:** Estimate $Y|X$ or $\mathbb{E}[Y|X]$ directly without resorting to an explicit parametric model.

Non Parametric Conditional Estimation

- Two heuristics:
  - $Y|X$ (or $\mathbb{E}[Y|X]$) is almost constant (or simple) in a neighborhood of $X$. (Kernel methods)
  - $Y|X$ (or $\mathbb{E}[Y|X]$) can be approximated by a model whose dimension depends on the complexity and the number of observation. (Quite similar to parametric model plus model selection...)

- Focus on **kernel methods**!
**Kernel Methods**

- **Idea:** The behavior of $Y|X$ is locally *constant* or simple!

**Kernel**

- Choose a kernel $K$ (think of a weighted neighborhood).
- For each $\tilde{X}$, compute a simple localized estimate of $Y|X$.
- Use this local estimate to take the decision.

- In regression, estimation of $\mathbb{E}[Y|X]$ is sufficient.
Example: $k$ Nearest-Neighbors (with $k = 3$)
Example: $k$ Nearest-Neighbors (with $k = 4$)
**k Nearest-Neighbors**

- Neighborhood $\mathcal{V}_x$ of $x$: $k$ learning samples closest from $x$.

### $k$-NN as local conditional density estimate

$$\mathbb{P}(\hat{Y} = 1|X) = \frac{\sum_{X_i \in \mathcal{V}_x} 1\{Y_i = +1\}}{|\mathcal{V}_x|}$$

- **KNN Classifier:**
  $$\hat{f}_{\text{KNN}}(X) = \begin{cases} 
+1 & \text{if } \mathbb{P}(\hat{Y} = 1|X) \geq \mathbb{P}(\hat{Y} = -1|X) \\
-1 & \text{otherwise}
\end{cases}$$

- **Lazy learning**: all the computations have to be done at prediction time.

- **Remark**: You can also use your favorite kernel estimator...
Example: KNN

k-NN with $k=1$

Decision region

Decision boundary

classes

Class1

Class2
Example: KNN

k-NN with k=5

Decision region

Decision boundary

classes
- Class1
- Class2

classes
- Class1
- Class2
Example: KNN

k-NN with $k=9$

Decision region

Decision boundary

- classes
  - Class1
  - Class2
Example: KNN

k-NN with $k=13$
Example: KNN

k-NN with \( k=17 \)

Decision region

Decision boundary

classes
- Class1
- Class2
Example: KNN

k-NN with $k=21$

Decision region:

- classes
  - Class1
  - Class2

Decision boundary:

- classes
  - Class1
  - Class2
Example: KNN
Example: KNN

k-NN with $k=29$

Decision region

Decision boundary

classes
- Class1
- Class2
Example: KNN

k-NN with k=33

Decision region

Decision boundary

classes
- Class1
- Class2
Example: KNN

k-NN with $k=37$

Decision region

Decision boundary

classes

- Class1
- Class2
Example: KNN

k-NN with $k=45$

Decision region

Decision boundary
Example: KNN

k-NN with $k=53$
Example: KNN

k-NN with k=61

Decision region

- PredictorA
- PredictorB
- Classes: Class1 (red), Class2 (blue)

Decision boundary

- PredictorA
- PredictorB
- Classes: Class1 (red), Class2 (blue)
Example: KNN

k-NN with $k=69$

Decision region

Decision boundary
Example: KNN
Example: KNN

k-NN with k=85
Example: KNN
Example: KNN

k-NN with k=109

Decision region

Decision boundary

classes
Class1
Class2

PredictorB
PredictorA
Example: KNN

k-NN with $k=117$

Decision region

Decision boundary

classes

Class 1

Class 2

classes

Class 1

Class 2
Example: KNN

k-NN with $k=125$
Example: KNN

k-NN with $k=133$

Decision region

Decision boundary

classes
- Class1
- Class2
Example: KNN

k-NN with $k=141$

Decision region

Decision boundary

classes

- Class1
- Class2
Example: KNN

k-NN with $k=149$

Decision region

Decision boundary

classes
- Class1
- Class2
Example: KNN

k-NN with $k=157$

Decision region

Decision boundary

classes

Class 1

Class 2
Example: KNN

k-NN with $k=165$
Example: KNN

k-NN with $k=173$

Decision region

Decision boundary
Example: KNN

k-NN with $k=181$

Decision region

Decision boundary

classes
- Class1
- Class2

PredictorA

PredictorB

0.2 0.4 0.6

0.2 0.4 0.6
Example: KNN

k-NN with k=189

Decision region

Decision boundary

classes
- Class1
- Class2
Example: KNN

k-NN with $k=197$

Decision region

Decision boundary

- classes
  - Class1
  - Class2
Regression and Local Averaging

A naive idea

- $\mathbb{E}[Y|X]$ can be approximated by a local average:

$$\hat{f}(X) = \frac{1}{|\{X_i \in \mathcal{N}(X)\}|} \sum_{X_i \in \mathcal{N}(X)} Y_i$$

where $\mathcal{B}(X)$ is a neighborhood of $X$.

- **Heuristic:**
  - If $X \rightarrow \mathbb{E}[Y|X]$ is regular then
    $$\mathbb{E}[Y|X] \simeq \mathbb{E}[\mathbb{E}[Y|X'] | X' \in \mathcal{N}(X)] = \mathbb{E}[Y|X' \in \mathcal{N}(X)]$$
  - Replace an expectation by an empirical average:
    $$\mathbb{E}[Y|X' \in \mathcal{N}(X)] \simeq \frac{1}{|\{X_i \in \mathcal{N}(X)\}|} \sum_{X_i \in \mathcal{N}(X)} Y_i$$
A Probabilistic Point of View

Regression and Local Averaging

Neighborhood and Size

- Most classical choice: \( \mathcal{N}(X) = \{ X', \|X - X'\| \leq h \} \) where \( \| . \| \) is a (pseudo) norm and \( h \) a size (bandwidth) parameter.
- In principle, the norm and \( h \) could vary with \( X \), and the norm can be replaced by a (pseudo) distance.
- Focus here on a fixed distance with a fixed bandwidth \( h \) case.

Bandwidth Heuristic

- A large bandwidth ensures that the average is taken on many samples and thus the variance is small...
- A small bandwidth is thus that the approximation \( \mathbb{E}[Y|X] \simeq \mathbb{E}[Y|X' \in \mathcal{N}(X)] \) is more accurate (small bias).
Weighted Local Averaging

Weighted Local Average

- Replace the neighborhood \( \mathcal{N}(X) \) by a decaying window function \( w(X, X') \).
- \( \mathbb{E}[Y|X] \) can be approximated by a weighted local average:

\[
\hat{f}(X) = \frac{\sum_i w(X, X') Y_i}{\sum_i w(X, X')}
\]

Kernel

- Most classical choice: \( w(X, X') = K \left( \frac{X-X'}{h} \right) \) where \( h \) the bandwidth is a scale parameter.
- Examples:
  - **Box kernel:** \( K(t) = 1_{\|t\| \leq 1} \) (Neighborhood)
  - **Triangular kernel:** \( K(t) = \max(1 - \|t\|, 0) \).
  - **Gaussian kernel:** \( K(t) = e^{-t^2/2} \)
- **Rk:** \( K \) and \( \lambda K \) yields the same estimate.
Nadaraya-Watson Heuristic

- Provided all the **densities** exist

\[ \mathbb{E}[Y|X] = \frac{\int Y p(X, Y) dY}{\int p(Y, X) dY} = \frac{\int Y p(X, Y) dY}{p(X)} \]

- Replace the unknown densities by their **estimates**:

\[ \hat{p}(X) = \frac{1}{n} \sum_{i=1}^{n} K(X - X_i) \]

\[ \hat{p}(X, Y) = \frac{1}{n} \sum_{i=1}^{n} K(X - X_i) K'(Y - Y_i) \]

- Now if \( K' \) is a kernel such that \( \int Y K'(Y) dY = 0 \) then

\[ \int Y \hat{p}(X, Y) dY = \frac{1}{n} \sum_{i=1}^{n} K(X - X_i) Y_i \]
From Density Estimation to Regression

Nadaraya-Watson

- Resulting estimator of $\mathbb{E}[Y|X]$
  \[
  \hat{f}(X) = \frac{\sum_{i=1}^{n} Y_i K_h(X - X_i)}{\sum_{i=1}^{n} K_h(X - X_i)}
  \]
- Same **local weighted average** estimator!

Bandwidth Choice

- Bandwidth $h$ of $K$ allows to **balance between bias and variance**.
- Theoretical analysis of the error is possible.
- The smoother the densities the easier the estimation but the optimal bandwidth depends on the unknown regularity!
Local Linear Estimation

Another Point of View on Kernel

- Nadaraya-Watson estimator:

\[
\hat{f}(X) = \frac{\sum_{i=1}^{n} Y_i K_h(X - X_i)}{\sum_{i=1}^{n} K_h(X - X_i)}
\]

- Can be viewed as a minimizer of

\[
\sum_{i=1}^{n} |Y_i - \beta|^2 K_h(X - X_i)
\]

- **Local regression** of order 0!

Local Linear Model

- Estimate \(E[Y|X]\) by \(\hat{f}(X) = \phi(X)^T \hat{\beta}(X)\) where \(\phi\) is any function of \(X\) and \(\hat{\beta}(X)\) is the minimizer of

\[
\sum_{i=1}^{n} |Y_i - \phi(X_i)^T \beta|^2 K_h(X - X_i).
\]
A Probabilistic Point of View

LOESS: LOcal polynomial regrESSion

1D Nonparametric Regression

- Assume that $X \in \mathbb{R}$ and let $\phi(X) = (1, X, \ldots, X^d)$.
- **LOESS estimate:** $\hat{f}(X) = \sum_{j=0}^{d} \hat{\beta}(X^{(j)}) X^j$ with $\hat{\beta}(X)$ minimizing

$$\sum_{i=1}^{n} |Y_i - \sum_{j=0}^{d} \beta^{(j)} X_i^j|^2 K_h(X - X_i).$$

- Most classical kernel used: Tricubic kernel

$$K(t) = \max(1 - |t|^3, 0)^3$$

- Most classical degree: 2...
- Local bandwidth choice such that a proportion of points belongs to the window.
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Fully Generative Modeling

**Idea:** If one knows the law of $(X, Y)$ everything is easy!

**Bayes formula**

With a slight abuse of notation,

$$
P(Y|X) = \frac{P((X, Y))}{P(X)}
= \frac{P(X|Y)P(Y)}{P(X)}
$$

**Generative Modeling:**

- Propose a model for $(X, Y)$ (or equivalently $X|Y$ and $Y$),
- Estimate it as a density estimation problem,
- Plug the estimate in the Bayes formula
- Plug the conditional estimate in the Bayes classifier.

**Rk:** Require to estimate $(X, Y)$ rather than only $Y|X$!

Great flexibility in the model design but may lead to complex computation.
Fully Generative Modeling

- Simpler setting in classification!

Bayes formula

\[
P(Y = k | X) = \frac{P(X | Y = k) P(Y = k)}{P(X)}
\]

- Binary Bayes classifier (the best solution)

\[
f^*(X) = \begin{cases} 
+1 & \text{if } P(Y = 1 | X) \geq P(Y = -1 | X) \\
-1 & \text{otherwise}
\end{cases}
\]

- **Heuristic**: Estimate those quantities and plug the estimations.
- By using different models/estimators for \(P(X | Y)\), we get different classifiers.
- **Rk**: No need to renormalize by \(P(X)\) to take the decision!
Discriminant Analysis (Gaussian model)

- The densities are modeled as multivariate normal, i.e.,
  \[ P(X \mid Y = k) \sim \mathcal{N}_{\mu_k, \Sigma_k} \]

- Discriminant functions:
  \[ g_k(X) = \ln(P(X \mid Y = k)) + \ln(P(Y = k)) \]
  \[ g_k(X) = -\frac{1}{2}(X - \mu_k)^\top \Sigma_k^{-1}(X - \mu_k) \]
  \[ -\frac{d}{2} \ln(2\pi) - \frac{1}{2} \ln(|\Sigma_k|) + \ln(P(Y = k)) \]

- QDA (different \( \Sigma_k \) in each class) and LDA (\( \Sigma_k = \Sigma \) for all \( k \))

- Beware: this model can be false but the methodology remains valid!
Discriminant Analysis

- The probability densities are Gaussian
- The effect of any decision rule is to divide the feature space into some decision regions $\mathcal{R}_1, \mathcal{R}_2$
- The regions are separated by decision boundaries
Discriminant Analysis

Quadratic Discriminant Analysis

- The probability densities are Gaussian
- The effect of any decision rule is to divide the feature space into some decision regions $R_1, R_2, \ldots, R_c$
- The regions are separated by decision boundaries
## Discriminant Analysis

### Estimation

In practice, we will need to estimate $\mu_k$, $\Sigma_k$ and $P_k := P(Y = k)$

- The estimate proportion $P(Y = k) = \frac{n_k}{n} = \frac{1}{n} \sum_{i=1}^{n} 1\{Y_i=k\}$
- Maximum likelihood estimate of $\hat{\mu}_k$ and $\hat{\Sigma}_k$ (explicit formulas)

- DA classifier

\[
\hat{f}_G(X) = \begin{cases} 
+1 & \text{if } \hat{g}_+1(X) \geq \hat{g}_{-1}(X) \\
-1 & \text{otherwise} 
\end{cases}
\]

- Decision boundaries: quadratic = degree 2 polynomials.
- If one imposes $\Sigma_{-1} = \Sigma_1 = \Sigma$ then the decision boundaries is a linear hyperplane.
Linear Discriminant Analysis

- $\sum \omega_1 = \sum \omega_2 = \Sigma$
- The decision boundaries are linear hyperplanes
Discriminant Analysis

Arbitrary Gaussian distributions lead to Bayes decision boundaries that are general quadratics.

- $\Sigma_{\omega_1} \neq \Sigma_{\omega_2}$
- Arbitary Gaussian distributions lead to Bayes decision boundaries that are general quadratics.
Example: LDA

Linear Discriminant Analysis

Decision region

Decision boundary

classes
Class1
Class2
Example: QDA

Quadratic Discriminant Analysis

Decision region

Decision boundary
Naive Bayes

- Classical algorithm using a crude modeling for $\mathbb{P}(X|Y)$:
  - Feature independence assumption:
    \[ \mathbb{P}(X|Y) = \prod_{l=1}^{d} \mathbb{P}(X^{(l)}|Y) \]
  - Simple featurewise model: binomial if binary, multinomial if finite and Gaussian if continuous

- If all features are continuous, similar to the previous Gaussian but with a diagonal covariance matrix!

- Very simple learning even in very high dimension!
Example: Naive Bayes
Naive Bayes with Density Estimation
Example: Naive Bayes

Naive Bayes with kernel density estimates

Decision region

Decision boundary

classes
Class1
Class2

classes
Class1
Class2
Other Models

- Other models of the world!

**Bayesian Approach**
- Generative Model plus prior on the parameters
- Inference thanks to the Bayes formula

**Graphical Models**
- Markov type models on Graphs

**Gaussian Processes**
- Multivariate Gaussian models

...
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Probabilistic and Optimization Framework

How to find a good function $f$ with a small risk

$$R(f) = \mathbb{E}[\ell(Y, f(X))]$$

**Canonical approach:** $\hat{f}_S = \arg\min_{f \in S} \frac{1}{n} \sum_{i=1}^{n} \ell(Y_i, f(X_i))$

**Problems**

- How to choose $S$?
- How to compute the minimization?

**A Probabilistic Point of View**

**Solution:** For $X$, estimate $Y|X$ plug this estimate in the Bayes classifier:

(Generalized) Linear Models, Kernel methods, $k$-nn, Naive Bayes, Tree, Bagging...

**An Optimization Point of View**

**Solution:** If necessary replace the loss $\ell$ by an upper bound $\bar{\ell}$ and minimize the empirical loss: SVR, SVM, Neural Network, Tree, Boosting...
Three Classical Methods in a Nutshell

Penalized Logistic Regression

- Let $f_\theta(X) = X^T \beta + \beta^{(0)}$ with $\theta = (\beta, \beta^{(0)})$.
- Find $\hat{\theta} = \arg \min \frac{1}{n} \sum_{i=1}^{n} \log \left(1 + e^{-Y_i f_\theta(X_i)}\right) + \lambda \|\beta\|_1$
- Classify using $\text{sign}(f_{\hat{\theta}})$

Support Vector Machine

- Let $f_\theta(X) = X^T \beta + \beta^{(0)}$ with $\theta = (\beta, \beta^{(0)})$.
- Find $\hat{\theta} = \arg \min \frac{1}{n} \sum_{i=1}^{n} \max (1 - Y_i f_\theta(X_i), 0) + \lambda \|\beta\|_2^2$
- Classify using $\text{sign}(f_{\hat{\theta}})$
Deep Learning

- Let $f_\theta(X)$ with $f$ a feed forward neural network outputing two values with a softmax layer as a last layer.

- Optimize by gradient descent the cross-entropy $\frac{1}{n} \sum_{i=1}^{n} \log \left( f_\theta(X_i)^{(Y_i)} \right)$

- Classify using $\text{sign}(f_\theta)$

- Those three methods rely on a similar heuristic: the optimization point of view!
The best solution \( f^* \) is the one minimizing
\[
f^* = \arg \min R(f) = \arg \min \mathbb{E}[\ell(Y, f(X))]
\]

One restricts \( f \) to a subset of functions \( S = \{f_\theta, \theta \in \Theta\} \)

One replaces the minimization of the average loss by the minimization of the average empirical loss
\[
\hat{f} = \hat{f}_\theta = \arg \min_{f_\theta, \theta \in \Theta} \frac{1}{n} \sum_{i=1}^{n} \ell(Y_i, f_\theta(X_i))
\]

Intractable for the \( \ell^0/1 \) loss!
**Convexification Strategy**

**Risk Convexification**
- Replace the loss $\ell(Y, f_\theta(X))$ by a convex upperbound $\bar{\ell}(Y, f_\theta(X))$ (surrogate loss).
- Minimize the average of the surrogate empirical loss

$$\tilde{f} = f_\hat{\theta} = \arg\min_{f_\theta, \theta \in \Theta} \frac{1}{n} \sum_{i=1}^{n} \bar{\ell}(Y_i, f_\theta(X_i))$$

- Use $\hat{f} = \text{sign}(\tilde{f})$

- Much easier optimization.

**Instantiation**
- Logistic (Revisited)
- Support Vector Machine
- (Deep) Neural Network
- Boosting
Classification Loss and Convexification

Convexification

- Replace the loss $\ell^{0/1}(Y, f(X))$ by
  $$\tilde{\ell}(Y, f(X)) = l(Yf(X))$$
  with $l$ a convex function.

- **Further mild assumption:** $l$ is decreasing, differentiable at 0 and $l'(0) < 0$. 
Classification Loss and Convexification

Classical convexification

- **Logistic loss**: \( \tilde{\ell}(Y, f(X)) = \log_2(1 + e^{-Yf(X)}) \) (Logistic / NN)
- **Hinge loss**: \( \tilde{\ell}(Y, f(X)) = (1 - Yf(X))_+ \) (SVM)
- **Exponential loss**: \( \tilde{\ell}(Y, f(X)) = e^{-Yf(X)} \) (Boosting...)
Properties

The Target is the Bayes Classifier

- The minimizer of
  \[
  \mathbb{E}[\tilde{\ell}(Y, f(X))] = \mathbb{E}[\ell(Yf(X))]
  \]
  is the Bayes classifier \( f^* = \text{sign}(2\eta(X) - 1) \)

Control of the Excess Risk

- It exists a convex function \( \Psi \) such that
  \[
  \Psi \left( \mathbb{E}[\ell^{0/1}(Y, \text{sign}(f(X)))] - \mathbb{E}[\ell^{0/1}(Y, f^*(X))] \right) \\
  \leq \mathbb{E}[\tilde{\ell}(Y, f(X))] - \mathbb{E}[\tilde{\ell}(Y, f^*(X))]
  \]

- Theoretical guarantee!
Logistic Revisited

- Ideal solution:
  \[ \hat{f} = \arg\min_{f \in S} \frac{1}{n} \sum_{i=1}^{n} \ell^{0/1}(Y_i, f(X_i)) \]

Logistic regression

- Use \( f(X) = X^\top \beta + \beta^{(0)}. \)
- Use the logistic loss \( \ell(y, f) = \log_2(1 + e^{-yf}), \) i.e. the negative log-likelihood.

- Different vision than the statistician but same algorithm!
Logistic Revisited

Logistic

Decision region

Decision boundary

classes
Class1
Class2

classes
Class1
Class2
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Ideal Separable Case

- Linear classifier: \( \text{sign}(X^T \beta + \beta(0)) \)
- Separable case: \( \exists (\beta, \beta(0)), \forall i, Y_i(X_i^T \beta + \beta(0)) > 0! \)

How to choose \((\beta, \beta(0))\) so that the separation is maximal?

- Strict separation: \( \exists (\beta, \beta(0)), \forall i, Y_i(X_i^T \beta + \beta(0)) \geq 1 \)
- Distance between \( X^T \beta + \beta(0) = 1 \) and \( X^T \beta + \beta(0) = -1 \):
  \[
  \frac{1}{2} \| \beta \| \quad \text{for} \quad X^T \beta + \beta(0) = 1 \quad \text{and} \quad X^T \beta + \beta(0) = -1.
  \]
  \[
  \frac{1}{2} \| \beta \| \quad \text{for} \quad X^T \beta + \beta(0) = 1 \quad \text{and} \quad X^T \beta + \beta(0) = -1.
  \]
- Maximizing this distance is equivalent to minimizing \( \frac{1}{2} \| \beta \|^2 \).
Ideal Separable Case

Separable SVM

- Constrained optimization formulation:
  \[
  \min \frac{1}{2} \|\beta\|^2 \quad \text{with} \quad \forall i, \ Y_i (X_i^T \beta + \beta^{(0)}) \geq 1
  \]

- Quadratic Programming setting.
- Efficient solver available...
What about the non separable case?

**SVM relaxation**

- Relax the assumptions
  \[
  \forall i, Y_i(X_i^\top \beta + \beta^{(0)}) \geq 1 \quad \text{to} \quad \forall i, Y_i(X_i^\top \beta + \beta^{(0)}) \geq 1 - s_i
  \]
  with the **slack variables** \( s_i \geq 0 \)

- Keep those slack variables as small as possible by minimizing
  \[
  \frac{1}{2}\|\beta\|^2 + C \sum_{i=1}^{n} s_i
  \]
  where \( C > 0 \) is the **goodness-of-fit strength**
Non Separable Case

SVM

- Constrained optimization formulation:
  \[
  \min \frac{1}{2} \|\beta\|^2 + C \sum_{i=1}^{n} s_i \quad \text{with} \quad \begin{cases} 
    \forall i, \ Y_i(X_i^T \beta + \beta^{(0)}) \geq 1 - s_i \\
    \forall i, \ s_i \geq 0
  \end{cases}
  \]

- **Hinge Loss** reformulation:
  \[
  \min \frac{1}{2} \|\beta\|^2 + C \sum_{i=1}^{n} \max(0, 1 - Y_i(X_i^T \beta + \beta^{(0)}))
  \]

- Constrained convex optimization algorithms vs gradient descent algorithms.
SVM as a Penalized Convex Relaxation

- Convex relaxation:

\[
\begin{align*}
\text{argmin} & \quad \frac{1}{2} \| \beta \|^2 + C \sum_{i=1}^{n} \max(1 - Y_i (X_i^T \beta + \beta(0)), 0) \\
\end{align*}
\]

\[
= \text{argmin} \frac{1}{n} \sum_{i=1}^{n} \max(1 - Y_i (X_i^T \beta + \beta(0)), 0) + \frac{1}{Cn} \frac{1}{2} \| \beta \|^2
\]

- Prop: \( \ell^{0/1}(Y_i, \text{sign}(X_i^T \beta + \beta(0))) \leq \max(1 - Y_i (X_i^T \beta + \beta(0)), 0) \)

Penalized convex relaxation (Tikhonov!)

\[
\begin{align*}
\frac{1}{n} \sum_{i=1}^{n} \ell^{0/1}(Y_i, \text{sign}(X_i^T \beta + \beta(0))) \\
\leq \frac{1}{n} \sum_{i=1}^{n} \max(1 - Y_i (X_i^T \beta + \beta(0)), 0) + \frac{1}{Cn} \frac{1}{2} \| \beta \|^2
\end{align*}
\]
Optimization Point of View

Support Vector Machine

Decision region

Decision boundary

classes

Class1

Class2
Constrained Minimization

Goal:

\[ \min_{x} f(x) \]

with

\[ \begin{cases} h_j(x) = 0, & j = 1, \ldots, p \\ g_i(x) \leq 0, & i = 1, \ldots, q \end{cases} \]

or rather with \( \text{argmin} \)!

Different Setting

- \( f, h_j, g_i \) differentiable
- \( f \) convex, \( h_j \) affine and \( g_i \) convex.

Feasibility

- \( x \) is **feasible** if \( h_j(x) = 0 \) and \( g_i(x) \leq 0 \).
- **Rk:** The set of feasible points may be empty.
### Optimization Point of View

#### Constrained Minimization

- **Goal:**
  \[ p^* = \min_x f(x) \quad \text{with} \quad \begin{cases} 
  h_j(x) = 0, & j = 1, \ldots, p \\
  g_i(x) \leq 0, & i = 1, \ldots, q 
  \end{cases} \]

#### Lagrangian

- **Def:**
  \[ \mathcal{L}(x, \lambda, \mu) = f(x) + \sum_{j=1}^{p} \lambda_j h_j(x) + \sum_{i=1}^{q} \mu_i g_i(x) \]

  with \( \lambda \in \mathbb{R}^p \) and \( \mu \in (\mathbb{R}^+)^q \).

- The \( \lambda_j \) and \( \mu_i \) are called the dual (or Lagrange) variables.

- **Prop:**
  \[ \max_{\lambda \in \mathbb{R}^p, \mu \in (\mathbb{R}^+)^q} \mathcal{L}(x, \lambda, \mu) = \begin{cases} 
  f(x) & \text{if } x \text{ is feasible} \\
  +\infty & \text{otherwise} 
  \end{cases} \]

  \[ \min_x \max_{\lambda \in \mathbb{R}^p, \mu \in (\mathbb{R}^+)^q} \mathcal{L}(x, \lambda, \mu) = p^* \]
Lagrangian

- **Def:**

\[ \mathcal{L}(x, \lambda, \mu) = f(x) + \sum_{j=1}^{p} \lambda_j h_j(x) + \sum_{i=1}^{q} \mu_i g_i(x) \]

with \( \lambda \in \mathbb{R}^p \) and \( \mu \in (\mathbb{R}^+)^q \).

Lagrangian Dual

- **Lagrangian dual function:**

\[ Q(\lambda, \mu) = \min_{x} \mathcal{L}(x, \lambda, \mu) \]

- **Prop:**

\[ Q(\lambda, \mu) \leq f(x), \text{ for all feasible } x \]

\[ \max_{\lambda \in \mathbb{R}^p, \mu \in (\mathbb{R}^+)^q} Q(\lambda, \mu) \leq \min_{x \text{ feasible}} f(x) \]
## Duality

**Primal**

- **Primal:**

\[
p^* = \min_{x \in \mathcal{X}} f(x) \text{ with } \begin{cases} 
  h_j(x) = 0, & j = 1, \ldots, p \\
  g_i(x) \leq 0, & i = 1, \ldots, q
\end{cases}
\]

**Dual**

- **Dual:**

\[
q^* = \max_{\lambda \in \mathbb{R}^p, \mu \in (\mathbb{R}^+)^q} Q(\lambda, \mu) = \max_{\lambda \in \mathbb{R}^p, \mu \in (\mathbb{R}^+)^q} \min_x L(x, \lambda, \mu)
\]

**Duality**

- **Always weak duality:**

\[
q^* \leq p^*
\]

\[
\max_{\lambda \in \mathbb{R}^p, \mu \in (\mathbb{R}^+)^q} \min_x L(x, \lambda, \mu) \leq \min_x \max_{\lambda \in \mathbb{R}^p, \mu \in (\mathbb{R}^+)^q} L(x, \lambda, \mu)
\]

- **Not always strong duality** \(q^* = p^*\).
Strong Duality

**Strong duality:**

\[
q^* = p^*
\]

\[
\max_{\lambda \in \mathbb{R}^p, \mu \in (\mathbb{R}^+)^q} \min_x \mathcal{L}(x, \lambda, \mu) = \min_x \max_{\lambda \in \mathbb{R}^p, \mu \in (\mathbb{R}^+)^q} \mathcal{L}(x, \lambda, \mu)
\]

- Allow to compute the solution of one problem from the other.
- Requires some assumptions!

**Strong Duality under Convexity and Slater’s Condition**

- \(f\) convex, \(h_j\) affine and \(g_i\) convex.
- **Slater’s condition:** it exists a feasible point such that \(h_j(x) = 0\) for all \(j\) and \(g_i(x) < 0\) for all \(i\).
- Sufficient to prove **strong duality**.
- **Rk:** If the \(g_i\) are affine, it suffices to have \(h_j(x) = 0\) for all \(j\) and \(g_i(x) \leq 0\) for all \(i\).
Karush-Kuhn-Tucker (KKT) Condition

- **Stationarity:**
  \[ \nabla_x \mathcal{L}(x^*, \lambda, \mu) = \nabla f(x^*) + \sum_j \lambda_j \nabla h_j(x^*) + \sum_i \mu_i \nabla g_i(x^*) = 0 \]

- **Primal admissibility:**
  \[ h_j(x^*) = 0 \quad \text{and} \quad g_i(x^*) \leq 0 \]

- **Dual admissibility:**
  \[ \mu_i \geq 0 \]

- **Complementary slackness:**
  \[ \mu_i g_i(x^*) = 0 \]

KKT Theorem

- If \( f \) convex, \( h_j \) affine and \( g_i \) convex, all are differentiable and strong duality holds then \( x^* \) is a solution of the primal problem if and only if the KKT condition holds.
SVM and Lagrangian

SVM

- Constrained optimization formulation:

\[
\min \frac{1}{2} \| \beta \|^2 + C \sum_{i=1}^{n} s_i \quad \text{with} \quad \begin{cases} 
\forall i, \, Y_i (X_i^\top \beta + \beta(0)) \geq 1 - s_i \\
\forall i, \, s_i \geq 0
\end{cases}
\]

SVM Lagrangian

- Lagrangian:

\[
\mathcal{L}(\beta, \beta(0), s, \alpha, \mu) = \frac{1}{2} \| \beta \|^2 + C \sum_{i=1}^{n} s_i + \sum_{i} \alpha_i (1 - s_i - Y_i (X_i^\top \beta + \beta(0))) - \sum_{i} \mu_i s_i
\]
SVM and KKT

KKT Optimality Conditions

- Stationarity:
  \[ \nabla_\beta L(\beta, \beta^{(0)}, s, \alpha, \mu) = \beta - \sum_i \alpha_i Y_i X_i = 0 \]
  \[ \nabla_{\beta^{(0)}} L(\beta, \beta^{(0)}, s, \alpha, \mu) = -\sum_i \alpha_i = 0 \]
  \[ \nabla_{s_i} L(\beta, \beta^{(0)}, s, \alpha, \mu) = C - \alpha_i - \mu_i = 0 \]

- Primal and dual admissibility:
  \[(1 - s_i - Y_i(X_i^T \beta + \beta^{(0)})) \leq 0, \quad s_i \geq 0, \quad \alpha_i \geq 0, \quad \text{and} \quad \mu_i \geq 0\]

- Complementary slackness:
  \[\alpha_i (1 - s_i - Y_i(X_i^T \beta + \beta^{(0)})) = 0 \quad \text{and} \quad \mu_i s_i = 0\]

Consequence

- \[\beta^* = \sum_i \alpha_i Y_i X_i\] and \[0 \leq \alpha_i \leq C\].
- If \[\alpha_i \neq 0\], \[X_i\] is called a support vector and either
  - \[s_i = 0\] and \[Y_i(X_i^T \beta^* + \beta^{(0)*}) = 1\] (margin hyperplane),
  - or \[\alpha_i = C\] (outliers).
- \[\beta^{(0)*} = Y_i - X_i^T \beta^*\] for any support vector with \[0 < \alpha_i < C\].
SVM Dual

SVM Lagrangian Dual
- Lagrangian Dual:
  \[ Q(\alpha, \mu) = \min_{\beta, \beta^{(0)}, s} L(\beta, \beta^{(0)}, s, \alpha, \mu) \]
- Prop:
  - if \( \sum_i \alpha_i Y_i \neq 0 \) or \( \exists i, \alpha_i + \mu_i \neq C \),
    \[ Q(\alpha, \mu) = -\infty \]
  - if \( \sum_i \alpha_i Y_i = 0 \) and \( \forall i, \alpha_i + \mu_i = C \),
    \[ Q(\alpha, \mu) = \sum_i \alpha_i - \frac{1}{2} \sum_{i,j} \alpha_i \alpha_j Y_i Y_j X_i^\top X_j \]

SVM Dual problem
- Dual problem is a Quadratic Programming problem:
  \[ \max_{0 \leq \alpha \leq C} Q(\alpha, \mu) \Leftrightarrow \max_{\alpha \geq 0, \mu \geq 0} \sum_i \alpha_i - \frac{1}{2} \sum_{i,j} \alpha_i \alpha_j Y_i Y_j X_i^\top X_j \]
- Involves the \( X_i \) only through their scalar products.
Mercer Theorem

Mercer Representation Theorem

- For any loss $\bar{\ell}$ and any increasing function $\Phi$, the minimizer in $\beta$ of
  $$\sum_{i=1}^{n} \bar{\ell}(Y_i, X_i^\top \beta + \beta^{(0)}) + \Phi(\|\beta\|_2)$$
  is a linear combination of the input points $\beta^* = \sum_{i=1}^{n} \alpha_i' X_i$.

- Minimization problem in $\alpha'$:
  $$\sum_{i=1}^{n} \bar{\ell}(Y_i, \sum_{j} \alpha_j' X_i^\top X_j + \beta^{(0)}) + \Phi(\|\beta\|_2)$$
  involving only the scalar product of the data.

- Optimal predictor requires only to compute scalar products.
  $\hat{f}^*(X) = X^\top \beta^* + \beta^{(0)},* = \sum_{i} \alpha_i' X_i^\top X$

- Transform a problem in dimension $\text{dim}(\mathcal{X})$ in a problem in dimension $n$.
- Direct minimization in $\beta$ can be more efficient...
Feature Engineering

- Art of creating **new features** from the existing one $X$.
- Example: add monomials $(X^{(i)})^2, X^{(i)}X^{(i')}...$
- Adding feature increases the dimension.

Feature Map

- Application $\phi : X \rightarrow \mathbb{H}$ with $\mathbb{H}$ an Hilbert space.
- Linear decision boundary in $\mathbb{H}$: $\phi(X) \top \beta + \beta^{(0)} = 0$ is **not an hyperplane anymore** in $X$.

**Heuristic**: Increasing dimension allows to make data almost linearly separable.
Polynomial Mapping

Polynomial Mapping of order 2

- \( \phi : \mathbb{R}^2 \to \mathbb{R}^6 \)

\[
\phi(X) = \left( (X^{(1)})^2, (X^{(2)})^2, \sqrt{2}X^{(1)}X^{(2)}, \sqrt{2}X^{(1)}, \sqrt{2}X^{(2)}, 1 \right)
\]

- Allow to solve the XOR classification problem with the hyperplane \( X^{(1)}X^{(2)} = 0 \).

Polynomial Mapping and Scalar Product

- **Prop:**

\[
\phi(X)^T \phi(X') = (1 + X^T X')^2
\]
SVM Primal and Dual

Primal, Lagrangian and Dual

- **Primal:**
  
  \[
  \min \|\beta\|^2 + C \sum_{i=1}^{n} s_i \quad \text{with} \quad \begin{cases} 
  \forall i, Y_i (\phi(X_i)^\top \beta + \beta(0)) \geq 1 - s_i \\
  \forall i, s_i \geq 0 
  \end{cases}
  \]

- **Lagrangian:**
  
  \[
  \mathcal{L}(\beta, \beta^{(0)}, s, \alpha, \mu) = \frac{1}{2} \|\beta\|^2 + C \sum_{i=1}^{n} s_i \\
  + \sum_{i} \alpha_i (1 - s_i - Y_i (\phi(X_i)^\top \beta + \beta^{(0)})) - \sum_{i} \mu_i s_i
  \]

- **Dual:**
  
  \[
  \max_{\alpha \geq 0, \mu \geq 0} Q(\alpha, \mu) \iff \max_{0 \leq \alpha \leq C} \sum_{i} \alpha_i - \frac{1}{2} \sum_{i,j} \alpha_i \alpha_j Y_i Y_j \phi(X_i)^\top \phi(X_j)
  \]

- **Optimal**
  
  \[
  \phi(X)^\top \beta^* + \beta^{(0),*} = \sum_i \alpha_i Y_i \phi(X)^\top \phi(X_i)
  \]

- **Only need to know to compute** \(\phi(X)^\top \phi(X')\) **to obtain the solution.**
Many algorithms (e.g. SVM) require only to be able to compute the scalar product $\phi(X)^\top \phi(X')$.

**Kernel**

- Any application

$$k : \mathcal{X} \times \mathcal{X} \to \mathbb{R}$$

is called a **kernel** over $\mathcal{X}$.

**Kernel Trick**

- Computing directly the kernel $k(x, x') = \phi(X)^\top \phi(X')$ may be easier than computing $\phi(X), \phi(X')$ and then the scalar product.

Here $k$ is defined from $\phi$.

Under some assumption on $k$, $\phi$ can be implicitly **defined** from $k$!
PDS Kernel

Positive Definite Symmetric Kernels

- A kernel $k$ is PDS if and only if
  - $k$ is symmetric, i.e.
    
    $$k(X, X') = k(X', X)$$
  - for any $N \in \mathbb{N}$ and any $(X_1, \ldots, X_N) \in X^N$,
    
    $$K = [k(X_i, X_j)]_{1 \leq i, j \leq N}$$
  
  is positive semi-definite, i.e. $\forall u \in \mathbb{R}^N$
    
    $$u^T Ku = \sum_{1 \leq i, j \leq N} u^{(i)} u^{(j)} k(X_i, X_j) \geq 0$$
  
  or equivalently all the eigenvalues of $K$ are non-negative.

- The matrix $K$ is called the **Gram matrix** associated to $(X_1, \ldots, X_N)$. 
Reproducing Kernel Hilbert Space

Moore-Aronsajn Theorem

- For any PDS kernel \( k : \mathcal{X} \times \mathcal{X} \rightarrow \mathbb{R} \), it exists a Hilbert space \( \mathbb{H} \subset \mathbb{R}^\mathcal{X} \) with a scalar product \( \langle \cdot, \cdot \rangle_\mathbb{H} \) such that
  - it exists a mapping \( \phi : \mathcal{X} \rightarrow \mathbb{H} \) satisfying
    \[
    k(X, X') = \langle \phi(X), \phi(X') \rangle_\mathbb{H}
    \]
  - the reproducing property holds, i.e. for any \( h \in \mathbb{H} \) and any \( X \in \mathcal{X} \)
    \[
    h(X) = \langle h, k(X, \cdot) \rangle_\mathbb{H}.
    \]

- By def., \( \mathbb{H} \) is a reproducing kernel Hilbert space (RKHS).
- \( \mathbb{H} \) is called the feature space associated to \( k \) and \( \phi \) the feature mapping.
- No unicity in general.
- \( \mathbf{Rk} \): if \( k(X, X') = \phi'(X)^\top \phi'(X') \) with \( \phi' : \mathcal{X} \rightarrow \mathbb{R}^p \) then
  - \( \mathbb{H} \) can be chosen as \( \{ X \mapsto \phi'(X)^\top \beta, \beta \in \mathbb{R}^p \} \) and \( \| X \mapsto \phi'(X)^\top \beta \|^2_\mathbb{H} = \| \beta \|^2_2 \).
  - \( \phi(X)(X') = X^\top X' \).
## Kernel Construction Machinery

### Separable Kernel
- For any function $\Psi : \mathcal{X} \to \mathbb{R}$, $k(X, X') = \Psi(X)\Psi(X')$ is PDS.

### Kernel Stability
- For any PDS kernels $k_1$ and $k_2$, $k_1 + k_2$ and $k_1k_2$ are PDS kernels.
- For any sequence of PDS kernels $k_n$ converging pointwise to a kernel $k$, $k$ is a PDS kernel.
- For any PDS kernel $k$ such that $|k| \leq r$ and any power series $\sum_n a_nz^n$ with $a_n \geq 0$ and a convergence radius larger than $r$, $\sum_n a_nk^n$ is a PDS kernel.
- For any PDS kernel $k$, the renormalized kernel $k'(X, X') = \frac{k(X, X')}{\sqrt{k(X, X)k(X', X')}}$ is a PDS kernel.
- Cauchy-Schwartz for PDS: $k(X, X')^2 \leq k(X, X)k(X', X')$
# Classical Kernels

## PDS Kernels

- **Vanilla kernel:**
  
  \[ k(X, X') = X^T X' \]

- **Polynomial kernel:**
  
  \[ k(X, X') = (1 + X^T X')^k \]

- **Gaussian RBF kernel:**
  
  \[ k(X, X') = \exp \left( -\gamma \|X - X'\|^2 \right) \]

- **Tanh kernel:**
  
  \[ k(X, X') = \tanh(aX^T X' + b) \]

- Most classical is the Gaussian RBF kernel...
- Lots of freedom to construct kernel for non classical data.
Representor Theorem

Let $k$ be a PDS kernel and $\mathbb{H}$ its corresponding RKHS, for any increasing function $\Phi$ and any function $L : \mathbb{R}^n \to \mathbb{R}$, the optimization problem

$$\arg\min_{h \in \mathbb{H}} L(h(X_1), \ldots, h(X_n)) + \Phi(\|h\|)$$

admits only solutions of the form

$$\sum_{i=1}^{n} \alpha'_i k(X_i, \cdot).$$

Examples:

- (kernelized) SVM
- (kernelized) Penalized Logistic Regression (Ridge)
- (kernelized) Penalized Regression (Ridge)
Kernelized SVM

**Primal**

- Constrained Optimization:
  \[
  \min_{f \in \mathbb{H}, \beta(0), s} \|f\|_\mathbb{H}^2 + C \sum_{i=1}^{n} s_i \quad \text{with} \quad \begin{cases} \forall i, Y_i(f(X_i) + \beta(0)) \geq 1 - s_i \\ \forall i, s_i \geq 0 \end{cases}
  \]
- Hinge loss:
  \[
  \min_{f \in \mathbb{H}, \beta(0)} \|f\|_\mathbb{H}^2 + C \sum_{i=1}^{n} \max(0, 1 - Y_i(f(X_i) + \beta(0)))
  \]
- Representer:
  \[
  \min_{\alpha', \beta(0)} \sum_{i,j} \alpha'_i \alpha'_j k(X_i, X_j) + C \sum_{i=1}^{n} \max(0, 1 - Y_i(\sum_j \alpha'_j k(X_j, X_i) + \beta(0)))
  \]

**Dual**

- Dual:
  \[
  \max_{\alpha \geq 0, \mu \geq 0} Q(\alpha, \mu) \iff \max_{0 \leq \alpha \leq C} \sum_{i} \alpha_i - \frac{1}{2} \sum_{i,j} \alpha_i \alpha_j Y_i Y_j k(X_i, X_j)
  \]
Support Vector Machine with polynomial kernel

Decision region

Decision boundary

classes

Class1

Class2
Support Vector Machine with Gaussian kernel

Decision region

Decision boundary

classes
- Class1
- Class2
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   - Hoeffding and Finite Class
   - McDiarmid and Rademacher Complexity
   - VC Dimension
   - Structural Risk Minimization

8 References
Simplified Models

Bias-Variance Issue

- Most complex models may not be the best ones due to the variability of the estimate.
- Naive idea: can we *simplify* our model without loosing too much?
  - by using only a subset of the variables?
  - by forcing the coefficients to be small?
- Can we do better than exploring all possibilities?
Linear Models

- **Setting**: Gen. linear model = prediction of $Y$ by $h(x^T \beta)$.

**Model coefficients**

- Model entirely specified by $\beta$.
- Coefficientwise:
  - $\beta^{(i)} = 0$ means that the $i$th covariate is not used.
  - $\beta^{(i)} \sim 0$ means that the $i$th covariate as a low influence...

- If some covariates are useless, better use a simpler model...

**Submodels**

- **Simplify** the model through a constraint on $\beta$!
- Examples:
  - Support: Impose that $\beta^{(i)} = 0$ for $i \notin I$.
  - Support size: Impose that $\|\beta\|_0 = \sum_{i=1}^d 1_{\beta^{(i)} \neq 0} < C$
  - Norm: Impose that $\|\beta\|_p < C$ with $1 \leq p$ (Often $p = 2$ or $p = 1$)
Sparsity

- \( \beta \) is sparse if its number of non-zero coefficients (\( \ell_0 \)) is small...
- Easy interpretation in terms of dimension/complexity.

Norm Constraint and Sparsity

- Sparsest solution obtained by definition with the \( \ell_0 \) norm.
- No induced sparsity with the \( \ell_2 \) norm...
- Sparsity with the \( \ell_1 \) norm (can even be proved to be the same as with the \( \ell_0 \) norm under some assumptions).
- Geometric explanation.
Constraint and Penalization

Constrained Optimization

- Choose a constant $C$.
- Compute $\beta$ as

$$\arg\min_{\beta \in \mathbb{R}^d, \|\beta\|_p \leq C} \frac{1}{n} \sum_{i=1}^{n} \bar{\ell}(Y_i, h(x_i^\top \beta))$$

Lagrangian Reformulation

- Choose $\lambda$ and compute $\beta$ as

$$\arg\min_{\beta \in \mathbb{R}^d} \frac{1}{n} \sum_{i=1}^{n} \bar{\ell}(Y_i, h(x_i^\top \beta)) + \lambda \|\beta\|_p'$$

with $p' = p$ except if $p = 0$ where $p' = 1$.
- Easier calibration... but no explicit model $S$.

**Rk:** $\|\beta\|_p$ is not scaling invariant if $p \neq 0$... 
- Initial rescaling issue.
Penalization

Penalized Linear Model
- Minimization of
  \[
  \arg\min_{\beta \in \mathbb{R}^d} \frac{1}{n} \sum_{i=1}^{n} \ell(Y_i, h(x_i^T \beta)) + \text{pen}(\beta)
  \]
  where \(\text{pen}(\beta)\) is a (sparsity promoting) penalty
- Variable selection if \(\beta\) is sparse.

Classical Penalties
- AIC: \(\text{pen}(\beta) = \lambda \|\beta\|_0\) (non-convex / sparsity)
- Ridge: \(\text{pen}(\beta) = \lambda \|\beta\|_2^2\) (convex / no sparsity)
- Lasso: \(\text{pen}(\beta) = \lambda \|\beta\|_1\) (convex / sparsity)
- Elastic net: \(\text{pen}(\beta) = \lambda_1 \|\beta\|_1 + \lambda_2 \|\beta\|_2^2\) (convex / sparsity)

- Easy optimization if \(\text{pen}\) (and the loss) is convex...
- **Need to specify \(\lambda\) to define a ML method!**
Penalized Gen. Linear Models

Classical Examples

- Penalized Least Squares
- Penalized Logistic Regression
- Penalized Maximum Likelihood
- SVM
- Tree pruning

- Sometimes used even if the parameterization is not linear...
Penalization and Cross-Validation

Practical Selection Methodology

- Choose a penalty family $\text{pen}_\lambda$.
- Compute a CV risk for the penalty $\text{pen}_\lambda$ for all $\lambda \in \Lambda$.
- Determine $\hat{\lambda}$ the $\lambda$ minimizing the CV risk.
- Compute the final model with the penalty $\text{pen}_{\hat{\lambda}}$.

- CV allows to select a ML method, penalized estimation with a penalty $\text{pen}_{\hat{\lambda}}$, not a single predictor hence the need of a final reestimation.

Why not using CV on a grid?

- Grid size scales exponentially with the dimension!
- **If the penalized minimization is easy**, much cheaper to compute the CV risk for all $\lambda \in \Lambda$...
- CV performs best when the set of candidates is not too big (or is structured...)

Optimization Point of View
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8. References
Perceptron (Rosenblatt 1957)

- Inspired from biology.
- Very simple (linear) model!
- Physical implementation and proof of concept.
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- Physical implementation and proof of concept.
Artificial Neuron and Logistic Regression

*Artificial Neuron*
- **Structure:**
  - Mix inputs with a *weighted sum*,
  - Apply a (non linear) *activation function* to this sum,
  - Possibly threshold the result to make a decision.
- Weights learned by minimizing a loss function.
- Equivalent to linear regression when using a linear activation function!

*Logistic unit*
- **Structure:**
  - Mix inputs with a *weighted sum*,
  - Apply the *logistic function* \( \sigma(t) = e^t / (1 + e^t) \),
  - Threshold at 1/2 to make a decision!
- Logistic weights learned by minimizing the -log-likelihood.
Multilayer Perceptron

MLP (Rumelhart, McClelland, Hinton - 1986)

- Multilayer Perceptron: cascade of layers of artificial neuron units.
- Optimization through a gradient descent algorithm with a clever implementation (Backprop).
- Construction of a function by composing simple units.
- MLP corresponds to a specific direct acyclic graph structure.
- Non convex optimization problem!
Multilayer Perceptron

Neural Network

Decision region

- Classes:
  - Class 1 (red)
  - Class 2 (blue)

Decision boundary

- Classes:
  - Class 1 (red)
  - Class 2 (blue)
Universal Approximation Theorem

Universal Approximation Theorem (Hornik, 1991)

- A **single hidden layer neural network** with a linear output unit can approximate any continuous function **arbitrarily well** given enough hidden units.

- Valid for most activation functions.
- No bounds on the number of required units... (Asymptotic flavor)
- A single hidden layer is sufficient but more may require less units.
Deep Neural Network

Deep Neural Network structure

- Deep cascade of layers!
- No conceptual novelty…
- But a **lot of tricks** allowing to obtain a good solution: clever initialization, better activation function, weight regularization, accelerated stochastic gradient descent, early stopping…
- Use of GPU and a lot of data…
- Very impressive results!
Deep Neural Network

Decision region

Decision boundary

H2O NN
Deep Learning

Family of Machine Learning algorithm combining:

- a (deep) multilayered structure,
- a clever optimization including initialization and regularization.

- Examples: Deep NN, AutoEncoder, Recursive NN, GAN, Transformer...
- Interpretation as a **Representation Learning**.
- **Transfer learning**: use as initialization a pretrained net.
- Very efficient and still evolving!
Le Net - Y. LeCun (1989)

- 6 hidden layer architecture.
- Drastic reduction of the number of parameters through a translation invariance principle (convolution).

- Required 3 days of training for 60 000 examples!
- Tremendous improvement.
- Representation learned through the task.

- Bigger and deeper layers and thus much more parameters.
- Clever initialization scheme, RELU, renormalization and use of GPU.
- 6 days of training for 1.2 millions images.
- Tremendous improvement...
Deep Convolutional Networks

Trends

- Bigger and bigger networks! (GoogLeNet / Residual Neural Network / Transformers...)
- More computational power to learn better representation.
- Work in Progress!
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Classification And Regression Trees

Tree principle (CART by Breiman (85) / ID3 by Quinlan (86))

- Construction of a recursive partition through a tree structured set of questions (splits around a given value of a variable)
- For a given partition, probabilistic approach and optimization approach yield the same predictor!
- A simple majority vote/averaging in each leaf

- Quality of the prediction depends on the tree (the partition).
- **Intuitively:**
  - small leaves lead to low bias, but large variance
  - large leaves lead to large bias, but low variance...
- **Issue:** Minim. of the (penalized) empirical risk is NP hard!
- Practical tree construction are all based on two steps:
  - a top-down step in which branches are created (branching)
  - a bottom-up in which branches are removed (pruning)
Branching

Greedy top-bottom approach

- Start from a single region containing all the data
- Recursively split those regions along a certain variable and a certain value

- **No regret strategy** on the choice of the splits!
- **Heuristic:** choose a split so that the two new regions are as *homogeneous* possible...
Branching

Optimization Point of View

Greedy top-bottom approach

- Start from a single region containing all the data
- Recursively split those regions along a certain variable and a certain value

- **No regret strategy** on the choice of the splits!
- **Heuristic:** choose a split so that the two new regions are as *homogeneous* possible...
Optimization Point of View

Branching

\[ X_1 < .5? \]
\[ X_2 < .7? \]

Greedy top-bottom approach

- Start from a single region containing all the data
- Recursively split those regions along a certain variable and a certain value

- **No regret strategy** on the choice of the splits!
- **Heuristic**: choose a split so that the two new regions are as *homogeneous* possible...
Branching

X_1 < .5?
Yes
No

X_1 < .2?
Yes
No

X_2 < .7?
Yes
No

Greedy top-bottom approach

- Start from a single region containing all the data
- Recursively split those regions along a certain variable and a certain value
- **No regret strategy** on the choice of the splits!
- **Heuristic**: choose a split so that the two new regions are as *homogeneous* possible...
Branching

Various definition of inhomogeneous

- **CART**: empirical loss based criterion (least squares/prediction error)
  \[ C(R, \bar{R}) = \sum_{x_i \in R} \bar{\ell}(y_i, y(R)) + \sum_{x_i \in \bar{R}} \bar{\ell}(y_i, y(\bar{R})) \]

- **CART**: Gini index (Classification)
  \[ C(R, \bar{R}) = \sum_{x_i \in R} p(R)(1 - p(R)) + \sum_{x_i \in \bar{R}} p(\bar{R})(1 - p(\bar{R})) \]

- **C4.5**: entropy based criterion (Information Theory)
  \[ C(R, \bar{R}) = \sum_{x_i \in R} H(R) + \sum_{x_i \in \bar{R}} H(\bar{R}) \]

- CART with Gini is probably the most used technique...
- Other criterion based on $\chi^2$ homogeneity or based on different local predictors (generalized linear models...
Branching

Choice of the split in a given region

- Compute the criterion for **all features and all possible splitting points**
  (necessarily among the data values in the region)
- Choose the split **minimizing** the criterion

**Variations:** split at all categories of a categorical variable using a clever category ordering (ID3), split at a fixed position (median/mean)

**Stopping rules:**
- when a leaf/region contains less than a prescribed number of observations
- when the region is sufficiently homogeneous...

- May lead to a quite complex tree: over-fitting possible!
- Additional pruning often use.
Pruning

- **Model selection** within the (rooted) subtrees of previous tree!
- Number of subtrees can be quite large, but the tree structure allows to find the best model efficiently.

**Key idea**
- The predictor in a leaf depends only on the values in this leaf.
- **Efficient bottom-up (dynamic programming) algorithm** if the criterion used satisfies an additive property

\[
C(T) = \sum_{\mathcal{L} \in T} c(\mathcal{L})
\]

- Example: AIC / CV.
Examples of criterion satisfying this assumptions

- AIC type criterion:
  \[
  \sum_{i=1}^{n} \bar{\ell}(y_i, f_L(x_i)(x_i)) + \lambda |T| = \sum_{L \in T} \left( \sum_{x_i \in L} \bar{\ell}(y_i, f_L(x_i)) + \lambda \right)
  \]

- Simple cross-Validation (with \((x'_i, y'_i)\) a different dataset):
  \[
  \sum_{i=1}^{n'} \bar{\ell}(y'_i, f_L(x'_i)) = \sum_{L \in T} \left( \sum_{x'_i \in L} \bar{\ell}(y'_i, f_L(x'_i)) \right)
  \]

- Limit over-fitting for a single tree.
- **Rk:** almost never used when combining several trees...
CART

Decision region

Decision boundary

classes

Class1

Class2
### CART: Pros and Cons

<table>
<thead>
<tr>
<th><strong>Pros</strong></th>
<th><strong>Cons</strong></th>
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<tr>
<td>- Leads to an easily interpretable model</td>
<td>- Greedy optimization</td>
</tr>
<tr>
<td>- Fast computation of the prediction</td>
<td>- Hard decision boundaries</td>
</tr>
<tr>
<td>- Easily deals with categorical features (and missing values)</td>
<td>- Lack of stability</td>
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</table>

**Pros**

- Leads to an easily interpretable model
- Fast computation of the prediction
- Easily deals with categorical features (and missing values)

**Cons**

- Greedy optimization
- Hard decision boundaries
- Lack of stability
Ensemble methods

- Lack of robustness for single trees.
- How to combine trees?

**Parallel construction**
- Construct several trees from bootstrapped samples and average the responses (**Bagging**)
- Add more randomness in the tree construction (**Random Forests**)

**Sequential construction**
- Construct a sequence of trees by reweighting sequentially the samples according to their difficulties (**AdaBoost**)
- Reinterpretation as a stagewise additive model (**Boosting**)

Optimization Point of View
Ensemble methods

Decision region

Bagging

Decision boundary

classes

Prediciton A

Prediciton B

Class 1

Class 2
Ensemble methods

Random Forest

Decision region

Decision boundary

classes
Class1
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AdaBoost

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classes

Class1

Class2
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Ensemble Methods

- **Averaging**: combine several models by averaging (bagging, random forests, ...)
- **Boosting**: construct a sequence of (weak) classifiers (XGBoost, Lightgbm)
- **Stacking**: use the outputs of several models as features (tpot, ...)

- Loss of interpretability but gain in performance
- Beware of overfitting with stacking: the second learning step should be done with fresh data.
- No end to end optimization as in deep learning!
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Empirical Risk Minimization

Empirical Risk Minimizer (ERM)

- For any loss $\ell$ and function class $S$, 
  \[ \hat{f} = \arg\min_{f \in S} \frac{1}{n} \sum_{i=1}^{n} \ell(Y_i, f(X_i)) = \arg\min_{f \in S} \mathcal{R}_n(f) \]

- Key property:
  \[ \mathcal{R}_n(\hat{f}) \leq \mathcal{R}_n(f), \forall f \in S \]

- **Minimization not always tractable in practice!**

- Focus on the $\ell^{0/1}$ case:
  - only algorithm is to try all the functions,
  - not feasible is there are many functions
  - but interesting hindsight!
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ERM and PAC Analysis

- Theoretical control of the random (error estimation) term:
  \[ R(\hat{f}) - R(f^*_S) \]

**Probably Almost Correct Analysis**

- **Theoretical guarantee** that
  \[ \mathbb{P}\left( R(\hat{f}) - R(f^*_S) \leq \epsilon_S(\delta) \right) \geq 1 - \delta \]
  for a suitable \( \epsilon_S(\delta) \geq 0 \).

- Implies:
  - \[ \mathbb{P}\left( R(\hat{f}) - R(f^*) \leq R(f^*_S) - R(f^*) + \epsilon_S(\delta) \right) \geq 1 - \delta \]
  - \[ \mathbb{E}\left[ R(\hat{f}) - R(f^*_S) \right] \leq \int_{0}^{+\infty} \delta_S(\epsilon) \, d\epsilon \]

- The result should hold without any assumption on the law \( P \! \)
A General Decomposition

- By construction:
  \[
  R(\tilde{f}) - R(f^*_S) = R(\tilde{f}) - R_n(\tilde{f}) + R_n(\tilde{f}) - R_n(f^*_S) + R_n(f^*_S) - R(f^*_S) \\
  \leq R(\tilde{f}) - R_n(\tilde{f}) + R_n(f^*_S) - R(f^*_S) \\
  \leq (R(\tilde{f}) - R(f^*_S)) - (R_n(\tilde{f}) - R_n(f^*_S))
  \]

Four possible upperbounds

- \[
  R(\tilde{f}) - R(f^*_S) \leq \sup_{f \in S} ((R(f) - R(f^*_S)) - (R_n(f) - R_n(f^*_S)))
  \]
- \[
  R(\tilde{f}) - R(f^*_S) \leq \sup_{f \in S} (R(f) - R_n(f)) + (R_n(f^*_S) - R(f^*_S))
  \]
- \[
  R(\tilde{f}) - R(f^*_S) \leq \sup_{f \in S} (R(f) - R_n(f)) + \sup_{f \in S} (R_n(f) - R(f))
  \]
- \[
  R(\tilde{f}) - R(f^*_S) \leq 2 \sup_{f \in S} |R(f) - R_n(f)|
  \]

- Supremum of centered random variables!
- Key: Concentration of each variable...
By construction, for any $f' \in \mathcal{S}$,

$$\mathcal{R}(f') = \mathcal{R}_n(f') + (\mathcal{R}(f') - \mathcal{R}_n(f'))$$

A uniform upper bound for the risk

Simultaneously $\forall f' \in \mathcal{S}$,

$$\mathcal{R}(f') \leq \mathcal{R}_n(f') + \sup_{f \in \mathcal{S}} (\mathcal{R}(f) - \mathcal{R}_n(f))$$

Supremum of centered random variables!

**Key**: Concentration of each variable...

Can be interpreted as a justification of the ERM!
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Concentration of the Empirical Loss

* Empirical loss:

\[ \mathcal{R}_n(f) = \frac{1}{n} \sum_{i=1}^{n} \ell^{0/1}(Y_i, f(X_i)) \]

**Properties**

- \( \ell^{0/1}(Y_i, f(X_i)) \) are i.i.d. random variables in \([0, 1]\).

**Concentration**

\[
\begin{align*}
\mathbb{P}(\mathcal{R}(f) - \mathcal{R}_n(f) \leq \epsilon) &\geq 1 - e^{-2ne^2} \\
\mathbb{P}(\mathcal{R}_n(f) - \mathcal{R}(f) \leq \epsilon) &\geq 1 - e^{-2ne^2} \\
\mathbb{P}(|\mathcal{R}_n(f) - \mathcal{R}(f)| \leq \epsilon) &\geq 1 - 2e^{-2ne^2}
\end{align*}
\]

- Concentration of sum of bounded independent variables!
- Hoeffding theorem.
- Equiv. to \( \mathbb{P}\left(\mathcal{R}(f) - \mathcal{R}_n(f) \leq \sqrt{\log(1/\delta)/(2n)}\right) \geq 1 - \delta \)
**Theorem**

Let $Z_i$ be a sequence of ind. centered r.v. supported in $[a_i, b_i]$ then

$$\mathbb{P}\left(\sum_{i=1}^{n} Z_i \geq \epsilon\right) \leq e^{-\frac{2\epsilon^2}{\sum_{i=1}^{n} (b_i - a_i)^2}}$$

**Proof ingredients:**

- **Chernov bounds:**
  $$\mathbb{P}\left(\sum_{i=1}^{n} Z_i \geq \epsilon\right) \leq \frac{\mathbb{E}[e^{\lambda \sum_{i=1}^{n} Z_i}]}{e^{\lambda \epsilon}} \leq \prod_{i=1}^{n} \mathbb{E}[e^{\lambda Z_i}]$$

- **Exponential moment bounds:** $\mathbb{E}[e^{\lambda Z_i}] \leq e^{\frac{\lambda^2 (b_i - a_i)^2}{8}}$

- **Optimization in $\lambda$**

**Prop:**

$$\mathbb{E}[e^{\lambda \sum_{i=1}^{n} Z_i}] \leq e^{\frac{\lambda^2 \sum_{i=1}^{n} (b_i - a_i)^2}{8}}$$
Hoeffding Inequality

**Theorem**

- Let $Z_i$ be a sequence of independent centered random variables supported in $[a_i, b_i]$ then

$$\Pr \left( \sum_{i=1}^{n} Z_i \geq \epsilon \right) \leq e^{-\frac{2\epsilon^2}{\sum_{i=1}^{n} (b_i-a_i)^2}}$$

- $Z_i = \frac{1}{n} \left( \mathbb{E} \left[ \ell^{0/1}(Y, f(X)) \right] - \ell^{0/1}(Y_i, f(X_i)) \right)$
- $\mathbb{E}[Z_i] = 0$ and $Z_i \in \left[ \frac{1}{n} \left( \mathbb{E} \left[ \ell^{0/1}(Y, f(X)) \right] - 1 \right), \frac{1}{n} \mathbb{E} \left[ \ell^{0/1}(Y, f(X)) \right] \right]$
- Concentration:

$$\Pr(\mathcal{R}(f) - \mathcal{R}_n(f) \geq \epsilon) \leq e^{-2n\epsilon^2}$$

- By symmetry,

$$\Pr(\mathcal{R}_n(f) - \mathcal{R}(f) \geq \epsilon) \leq e^{-2n\epsilon^2}$$

- Combining the two yields

$$\Pr(|\mathcal{R}_n(f) - \mathcal{R}(f)| \geq \epsilon) \leq 2e^{-2n\epsilon^2}$$
Finite Class Case

Concentration

- If $S$ is finite of cardinality $|S|$, 

\[
P\left( \sup_f (\mathcal{R}(f) - \mathcal{R}_n(f)) \leq \sqrt{\frac{\log |S| + \log(1/\delta)}{2n}} \right) \geq 1 - \delta
\]

\[
P\left( \sup_f |\mathcal{R}_n(f) - \mathcal{R}(f)| \leq \sqrt{\frac{\log |S| + \log(1/\delta)}{2n}} \right) \geq 1 - 2\delta
\]

- Control of the supremum by a quantity depending on the cardinality and the probability parameter $\delta$.

- Simple combination of Hoeffding and a union bound.
Finite Class Case

PAC Bounds

- If $S$ is finite of cardinality $|S|$, with proba greater than $1 - 2\delta$
  \[
  R(\hat{f}) - R(f^*_S) \leq \sqrt{\frac{\log |S| + \log(1/\delta)}{2n}} + \sqrt{\frac{\log(1/\delta)}{2n}}
  \leq 2 \sqrt{\frac{\log |S| + \log(1/\delta)}{2n}}
  \]

- If $S$ is finite of cardinality $|S|$, with proba greater than $1 - \delta$, simultaneously $\forall f' \in S$,
  \[
  R(f') \leq R_n(f') + \sqrt{\frac{\log |S| + \log(1/\delta)}{2n}}
  \leq R_n(f') + \sqrt{\frac{\log |S|}{2n}} + \sqrt{\frac{\log(1/\delta)}{2n}}
  \]
Finite Class Case

PAC Bounds

- If $S$ is finite of cardinality $|S|$, with proba greater than $1 - 2\delta$
  \[ \mathcal{R}(\hat{f}) - \mathcal{R}(f^*_S) \leq \sqrt{\frac{\log |S|}{2n}} + \sqrt{\frac{2 \log(1/\delta)}{n}} \]

- If $S$ is finite of cardinality $|S|$, with proba greater than $1 - \delta$, simultaneously $\forall f' \in S$,
  \[ \mathcal{R}(f') \leq \mathcal{R}_n(f') + \sqrt{\frac{\log |S|}{2n}} + \sqrt{\frac{\log(1/\delta)}{2n}} \]

- Risk increases with the cardinality of $S$.
- Similar issue in cross-validation!
- No direct extension for an infinite $S$...
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Concentration of the Supremum of Empirical Losses

- Supremum of Empirical losses:
  \[ \Delta_n(S)(X_1, \ldots, X_n) = \sup_{f \in S} \mathcal{R}(f) - \mathcal{R}_n(f) \]
  \[= \sup_{f \in S} \left( \mathbb{E}[\ell^{0/1}(Y, f(X))] - \frac{1}{n} \sum_{i=1}^{n} \ell^{0/1}(Y_i, f(X_i)) \right) \]

Properties

- Bounded difference:
  \[ |\Delta_n(S)(X_1, \ldots, X_i, \ldots X_n) - \Delta_n(S)(X_1, \ldots X'_i, \ldots, X_n)| \leq 1/n \]

Concentration

\[ \mathbb{P}(\Delta_n(S) - \mathbb{E}[\Delta_n(S)] \leq \epsilon) \geq 1 - e^{-2n\epsilon^2} \]

- Concentration of bounded difference function.
- Generalization of Hoeffding theorem: McDiarmid Theorem.
McDiarmid Inequality

**Bounded difference function**

- $g : \mathcal{X}^n \rightarrow \mathbb{R}$ is a bounded difference function if it exist $c_i$ such that
  \[
  \forall (X_i)_{i=1}^n, (X'_i)_{i=1}^n \in \mathbb{R},
  \quad |g(X_1, \ldots, X_i, \ldots, X_n) - g(X_1, \ldots, X'_i, \ldots, X_n)| \leq c_i
  \]

**Theorem**

- If $g$ is a bounded difference function and $X_i$ are independent random variables then
  \[
  \mathbb{P}(g(X_1, \ldots, X_n) - \mathbb{E}[g(X_1, \ldots, X_n)] \geq \epsilon) \leq e^{\frac{-2\epsilon^2}{\sum_{i=1}^n c_i^2}}
  \]
  \[
  \mathbb{P}(\mathbb{E}[g(X_1, \ldots, X_n)] - g(X_1, \ldots, X_n) \geq \epsilon) \leq e^{\frac{-2\epsilon^2}{\sum_{i=1}^n c_i^2}}
  \]

- **Proof ingredients:**
  - Chernov bounds
  - Martingale decomposition...
McDiarmid Inequality

**Theorem**

- If $g$ is a bounded difference function and $X_i$ are independent random variables then

$$
P(g(X_1, \ldots, X_n) - \mathbb{E}[g(X_1, \ldots, X_n)] \geq \epsilon) \leq e^{\sum_{i=1}^{-2\epsilon^2} c_i^2}
$$

- Using $g = \Delta_n(S)$ for which $c_i = 1/n$ yields immediately

$$
P(\Delta_n(S) - \mathbb{E}[\Delta_n(S)] \geq \epsilon) \leq e^{\sum_{i=1}^{-2\epsilon^2} c_i^2} = e^{-2n\epsilon^2}
$$

- We derive then

$$
P(\Delta_n(S) \geq \mathbb{E}[\Delta_n(S)] + \epsilon) \leq e^{\sum_{i=1}^{-2\epsilon^2} c_i^2} = e^{-2n\epsilon^2}
$$

- It remains to upperbound

$$
\mathbb{E}[\Delta_n] = \mathbb{E}
\left[
\sup_{f \in S} \mathcal{R}(f) - \mathcal{R}_n(f)
\right]
$$
Rademacher Complexity

Theorem

- Let \( \sigma_i \) be a sequence of i.i.d. random symmetric Bernoulli variables (Rademacher variables):

\[
\mathbb{E} \left[ \sup_{f \in S} (\mathcal{R}(f) - \mathcal{R}_n(f)) \right] \leq 2 \mathbb{E} \left[ \sup_{f \in S} \frac{1}{n} \sum_{i=1}^{n} \sigma_i \ell^{0/1}(Y_i, f(X_i)) \right]
\]

Rademacher complexity

- Let \( B \subset \mathbb{R}^n \), the Rademacher complexity of \( B \) is defined as

\[
R_n(B) = \mathbb{E} \left[ \sup_{b \in B} \frac{1}{n} \sum_{i=1}^{n} \sigma_i b_i \right]
\]

- Theorem gives an upper bound of the expectation in terms of the average Rademacher complexity of the random set \( B_n(S) = \{(\ell^{0/1}(Y_i, f(X_i)))_{i=1}^{n}, f \in S\} \).

- Back to finite setting: This set is at most of cardinality \( 2^n \)!
Finite Set Rademacher Complexity Bound

**Theorem**

- If $B$ is finite and such that $\forall b \in B, \frac{1}{n} \| b \|_2^2 \leq M^2$, then

$$R_n(B) = \mathbb{E} \left[ \sup_{b \in B} \frac{1}{n} \sum_{i=1}^{n} \sigma_i b_i \right] \leq \sqrt{\frac{2M^2 \log |B|}{n}}$$

- If $B = B_n(S) = \{(\ell^0/1( Y_i, f(X_i)))_{i=1}^{n}, f \in S\}$, we have $M = 1$ and thus

$$R_n(B) \leq \sqrt{\frac{2 \log |B_n(S)|}{n}}$$

- We obtain immediately

$$\mathbb{E} \left[ \sup_{f \in S} (\mathcal{R}(f) - \mathcal{R}_n(f)) \right] \leq \mathbb{E} \left[ \sqrt{\frac{8 \log |B_n(S)|}{n}} \right].$$
Finite Set Rademacher Complexity Bound

Theorem

- With probability greater than $1 - 2\delta$,
  \[ \mathcal{R}(\hat{f}) - \mathcal{R}(f^*_S) \leq \mathbb{E} \left[ \sqrt{\frac{8 \log |B_n(S)|}{n}} \right] + \sqrt{\frac{2 \log(1/\delta)}{n}} \]

- With probability greater than $1 - \delta$, simultaneously $\forall f' \in S$
  \[ \mathcal{R}(f') \leq \mathcal{R}_n(f') + \mathbb{E} \left[ \sqrt{\frac{8 \log |B_n(S)|}{n}} \right] + \sqrt{\frac{\log(1/\delta)}{2n}} \]

- This is a direct consequence of the previous bound.
Corollary

- If $S$ is finite then with probability greater than $1 - 2\delta$

$$
\mathcal{R}(\hat{f}) - \mathcal{R}(f_S^*) \leq \sqrt{\frac{8 \log |S|}{n}} + \sqrt{\frac{2 \log(1/\delta)}{n}}
$$

- If $S$ is finite then with probability greater than $1 - \delta$, simultaneously $\forall f' \in S$

$$
\mathcal{R}(f') \leq \mathcal{R}_n(f') + \sqrt{\frac{8 \log |S|}{n}} + \sqrt{\frac{\log(1/\delta)}{2n}}
$$

- It suffices to notice that

$$
|B_n(S)| = |\{(\ell^{0/1}(Y_i, f(X_i)))_{i=1}^n, f \in S\}| \leq |S|
$$
Empirical Risk Minimization

Finite Set Rademacher Complexity Bound

- Same result with Hoeffding but with **better** constants!

\[
\mathcal{R}(\hat{f}) - \mathcal{R}(f^*_S) \leq \sqrt{\frac{\log |S|}{2n}} + \sqrt{\frac{2 \log(1/\delta)}{n}}
\]

\[
\mathcal{R}(f') \leq \mathcal{R}_n(f') + \sqrt{\frac{\log |S|}{2n}} + \sqrt{\frac{\log(1/\delta)}{2n}}
\]

- Difference due to the **crude** upperbound of

\[
\mathbb{E} \left[ \sup_{f \in S} (\mathcal{R}(f) - \mathcal{R}_n(f)) \right]
\]

- **Why bother?:** We do not have to assume that \( S \) is finite!

\[
|B_n(S)| \leq 2^n
\]
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Back to the Bound

**Theorem**

$$
\mathbb{E} \left[ \sup_{f \in S} (\mathcal{R}(f) - \mathcal{R}_n(f)) \right] \leq \mathbb{E} \left[ \sqrt{\frac{8 \log |B_n(S)|}{n}} \right]
$$

- **Key quantity:** \( \mathbb{E} \left[ \sqrt{\frac{8 \log |B_n(S)|}{n}} \right] \)

- Hard to control due to its structure!

**A first data dependent upperbound**

$$
\mathbb{E} \left[ \sqrt{\frac{8 \log |B_n(S)|}{n}} \right] \leq \sqrt{\frac{8 \log \mathbb{E}[|B_n(S)|]}{n}} \quad \text{(Jensen)}
$$

- Depends on the unknown \( P \)!
Shattering Coefficient

Shattering Coefficient (or Growth Function)

- The shattering coefficient of the class $S$, $s(S, n)$, is defined as
  $$s(S, n) = \sup \left\{ \left| \left\{ (f^0/1(Y_i, f(X_i)))_{i=1}^n \right\} \right| \right\}_{f \in S}$$
  $$\left((X_1, Y_1), \ldots, (X_n, Y_n)\right) \in (\mathcal{X} \times \{-1, 1\})^n$$

- By construction, $|B_n(S)| \leq s(S, n) \leq \min(2^n, |S|)!$

A data independent upperbound

$$\mathbb{E} \left[ \frac{8 \log |B_n(S)|}{n} \right] \leq \sqrt{\frac{8 \log s(S, n)}{n}}$$
Shattering Coefficient

Theorem

- With probability greater than $1 - 2\delta$,
  \[
  \mathcal{R}(\hat{f}) - \mathcal{R}(f^*_S) \leq \sqrt{\frac{8 \log s(S, n)}{n}} + \sqrt{\frac{2 \log(1/\delta)}{n}}
  \]

- With probability greater than $1 - \delta$, simultaneously $\forall f' \in S$,
  \[
  \mathcal{R}(f') \leq \mathcal{R}_n(f') + \sqrt{\frac{8 \log s(S, n)}{n}} + \sqrt{\frac{\log(1/\delta)}{2n}}
  \]

- Depends only on the class $S$!
**VC Dimension**

- The VC dimension $d_{VC}$ of $S$ is defined as the largest integer $d$ such that $s(S, d) = 2^d$
- The VC dimension can be infinite!

**VC Dimension and Dimension**

- **Prop:** If $\text{span}(S)$ corresponds to the sign of functions in a linear space of dimension $d$ then $d_{VC} \leq d$.
- VC dimension similar to the usual dimension.
VC Dimension and Sauer’s Lemma

Sauer’s Lemma

- If the VC dimension $d_{VC}$ of $S$ is finite

$$s(S, n) \leq \begin{cases} 2^n & \text{if } n \leq d_{VC} \\ \left(\frac{en}{d_{VC}}\right)^{d_{VC}} & \text{if } n > d_{VC} \end{cases}$$

- **Cor.**: $\log s(S, n) \leq d_{VC} \log \left(\frac{en}{d_{VC}}\right)$ if $n > d_{VC}$.
VC Dimension and PAC Bounds

PAC Bounds

- If $S$ is of VC dimension $d_{VC}$ then if $n > d_{VC}$
- With probability greater than $1 - 2\delta$,
  \[
  \mathcal{R}(\hat{f}) - \mathcal{R}(f^*_S) \leq \sqrt{\frac{8d_{VC} \log \left( \frac{en}{d_{VC}} \right)}{n}} + \sqrt{\frac{2 \log(1/\delta)}{n}}
  \]
- With probability greater than $1 - \delta$, simultaneously $\forall f' \in S$,
  \[
  \mathcal{R}(f') \leq \mathcal{R}_n(f') + \sqrt{\frac{8d_{VC} \log \left( \frac{en}{d_{VC}} \right)}{n}} + \sqrt{\frac{\log(1/\delta)}{2n}}
  \]

- **Rk:** If $d_{VC} = +\infty$ no uniform PAC bounds exists!
PAC Bounds

- Let $\pi_f > 0$ such that $\sum_{f \in S} \pi_f = 1$
- With proba greater than $1 - 2\delta$,
  \[ R(\hat{f}) - R(f^*_S) \leq \sqrt{\frac{\log(1/\pi_f)}{2n}} + \sqrt{\frac{2\log(1/\delta)}{n}} \]
- With proba greater than $1 - \delta$, simultaneously $\forall f' \in S$,
  \[ R(f') \leq R_n(f') + \sqrt{\frac{\log(1/\pi_f)}{2n}} + \sqrt{\frac{\log(1/\delta)}{2n}} \]

- Very similar proof than the uniform one!
- Much more interesting idea when combined with several models...
Assume we have a countable collection of set \((S_m)_{m \in \mathcal{M}}\) and let \(\pi_m\) be such that \(\sum_{m \in \mathcal{M}} \pi_m = 1\).

**Non Uniform Risk Bound**

With probability \(1 - \delta\), simultaneously for all \(m \in \mathcal{M}\) and all \(f \in S_m\),

\[
\mathcal{R}(f) \leq \mathcal{R}_n(f) + \mathbb{E} \left[ \sqrt{\frac{8 \log |B_n(S_m)|}{n}} \right] + \sqrt{\frac{\log(1/\pi_m)}{2n}} + \sqrt{\frac{\log(1/\delta)}{2n}}
\]

**Structural Risk Minimization**

Choose \(\hat{f}\) as the minimizer over \(m \in \mathcal{M}\) and \(f \in S_m\) of

\[
\mathcal{R}_n(f) + \mathbb{E} \left[ \sqrt{\frac{8 \log |B_n(S_m)|}{n}} \right] + \sqrt{\frac{\log(1/\pi_m)}{2n}}
\]

- Mimics the minimization of the integrated risk!
SRM and PAC Bound

PAC Bound

- If $\hat{f}$ is the SRM minimizer then with probability $1 - 2\delta$,

$$\mathcal{R}(\hat{f}) \leq \inf_{m \in \mathcal{M}} \inf_{f \in \mathcal{S}_m} \left( \mathcal{R}(f) + \mathbb{E} \left[ \sqrt{8 \log |B_n(S_m)|} \frac{\log(1/\pi_m)}{n} \right] + \sqrt{\log(1/\delta)} \right)$$

$$+ \sqrt{\frac{2 \log(1/\delta)}{n}}$$

- The SRM minimizer balances the risk $\mathcal{R}(f)$ and the upper bound on the estimation error $\mathbb{E} \left[ \sqrt{8 \log |B_n(S_m)|} \frac{\log(1/\pi_m)}{n} \right] + \sqrt{\frac{\log(1/\pi_m)}{2n}}$.

- $\mathbb{E} \left[ \sqrt{8 \log |B_n(S_m)|} \frac{\log(1/\pi_m)}{n} \right]$ can be replaced by an upper bound (for instance a VC based one)...
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