Introduction to Machine Learning

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Course Outline

- 1. Introduction
- 2. Preprocessing Data
- 3. Linear Regression
- 4. Classification Model: Logistic Regression
- How to Evaluate a Model
- 6. Tree Based Method
- 7. Unsupervised learning: Clustering Methods

In every chapter we will have some example in python

Prerequisites

Basic knowledge of:

- Statistic
- Probability
- Linear Algebra: matrix calculus
- **Calculus**: derivatives, gradient, ecc...

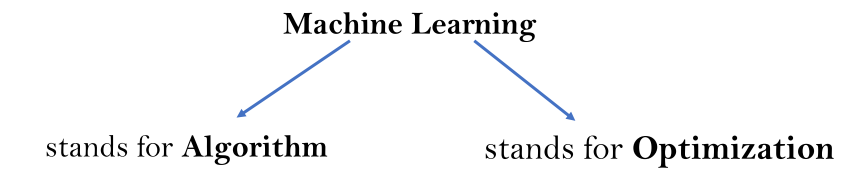
No prerequisites for programming

Recommended reading

- James, G., Witten, D., Hastie, T., & Tibshirani, R. (2013). *An introduction to statistical learning* (Vol. 112, p. 18). New York: springer.
- Hastie, T., Tibshirani, R., Friedman, J. H., & Friedman, J. H. (2009). The elements of statistical learning: data mining, inference, and prediction (Vol. 2, pp. 1-758). New York: springer.
- Rogers, S., & Girolami, M. (2016). A first course in machine learning. Chapman and Hall/CRC.
- Many resources available online, stackoverflow, standford and MIT courses etc...

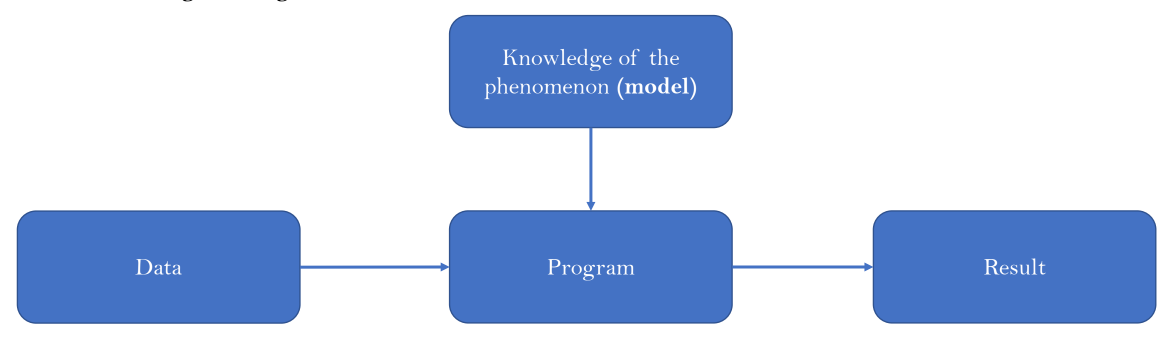
Chapter 1: Introduction

What is Machine Learning



A paradigm shift: from knowledge to data

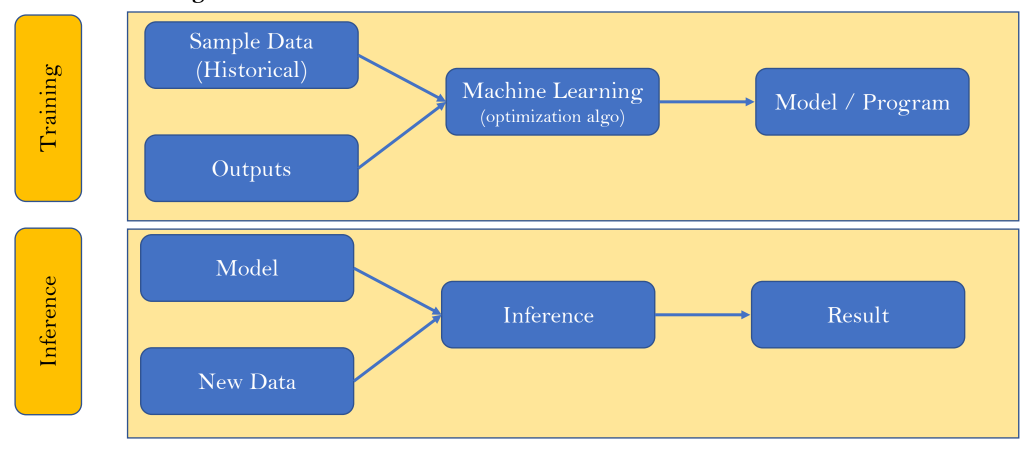
Traditional Programming:



Data are used to validate/test assumption of the pre-existing model

A paradigm shift: from knowledge to data

Machine Learning:



Data are used to build (train) a model used to get results (predictions) during inference \rightarrow where is the knowledge about the process? (see next slides)

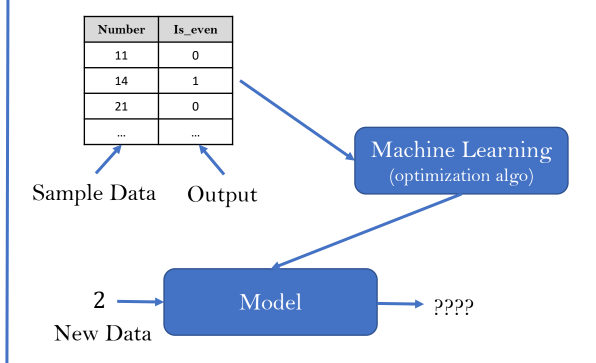
A paradigm shift: from knowledge to data

Write a program that output if a number is even or odd:

Traditional Programming:

A number is even if the rest of the division by 2 is 0

Machine Learning:

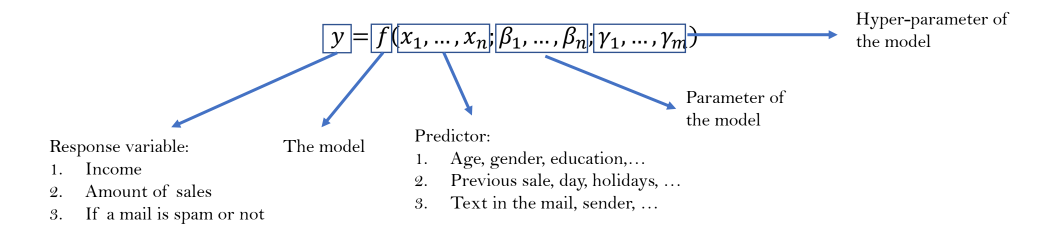


This approach will not work \rightarrow no 'simple' patter in the data

Machine Learning Foundamentals

Which Problems do ML solves?

ML Aims to learn (approximate) a model f, that relate the features x_i to the response y



Machine Learning Foundamentals

$$y = f(x_1, \dots, x_n; \beta_1, \dots, \beta_n; \gamma_1, \dots, \gamma_m)$$

How do we learn f?

The aim is to find "the best" model f given a dataset $D = \{Y_i, X_i\}$ i = 1, ..., n where n are the rows of D, X_i is the matrix of the predictor variables and Y_i is the response variable.

How to formalize the expression "the best"? → Optimization

We fix the shape for the model (e.g. linear model, tree based model, ecc..) and solve the following optimization problem:

$$\beta, \gamma = argmin_{\beta,\gamma} L(Y, f(X; \beta; \gamma))$$
 usually simplified* as $\beta = argmin_{\beta} L(Y, f(X; \beta; \gamma^*))$

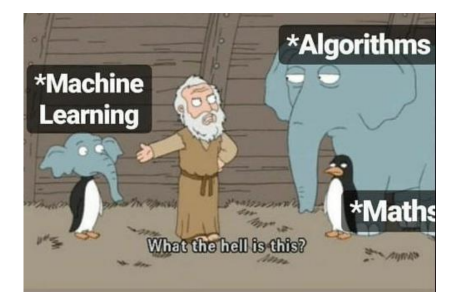
L is called the **Loss Function** and measures how well our model approximate the data

Machine Learning Foundamentals

$$y = f(x_1, ..., x_n; \beta_1, ..., \beta_n; \gamma_1, ..., \gamma_m)$$
How do we learn f ?
$$\beta = argmin_{\beta} L(Y, f(X; \beta; \gamma^*))$$
 (1)

In this course we will:

- Study several shape for f
- Understand the parameter & hyper-parameter of the f
- Define suitable loss function L for the problem at hand
- Understand how to compute hyperparameter
- Understand how to solve the minimization problem in (1)
- Understand the final result in term of business insight



Type of Problem

$$y = f(x_1, \dots, x_n)$$

Supervised

We observe both $x_i, y \rightarrow$ fit model that relate predictor to response

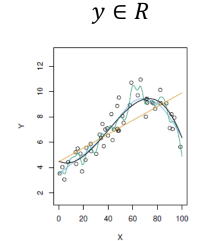
Unsupervised

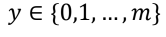
We observe only $x_i \rightarrow$ understand the relationships between the variables

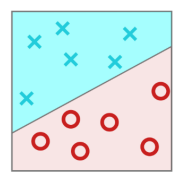
Regression

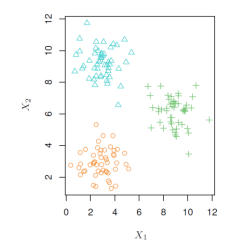
Classification

Clustering Methods

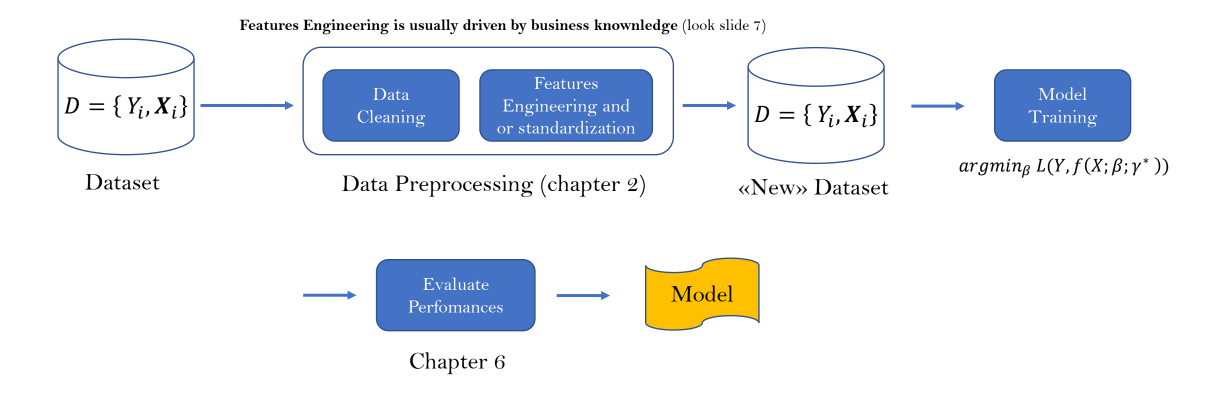








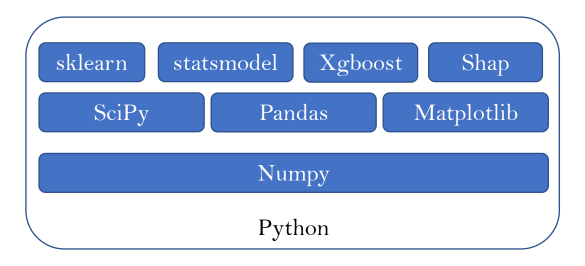
Machine Learning Pipeline



Python

Python is the most diffuse programming languages for ML applications

https://www.python.org/



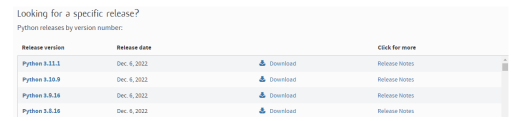
- **Numpy**: support for multidimensional array and mathematical function
- SciPy: scientific computing in python
- **Pandas**: data/table manipulation and analysis
- Matplotlib: Plotting graph
- Sklearn, statsmodel, (Xgboost): training of ML model
- **(Shap)**: variables explanation for complex model

Install python

Go to the website https://www.python.org/downloads/

Download the latest version of Python

Download Python 3.11.1



- I suggest to install version 3.10 or 3.9 (usually with the latest is more difficult to install package)
 - remember to add python to the \$PATH env variables (it will ask)
 - Install Jupyter notebook/lab at https://jupyter.org/install

Jupyter Notebook

Install the classic Jupyter Notebook with:

pip install notebook

To run the notebook:

jupyter notebook

pip is the packege manager of python

Pip install <package_name> == <package_version>
Pip uninstall <package_name>

```
Microsoft Windows [Versione 10.0.19044.2251]
(c) Microsoft Corporation. Tutti i diritti sono riservati.

C:\Users\LEI00068>pip install notebook

Prompt deicomandi

Microsoft Windows [Versione 10.0.19044.2251]
(c) Microsoft Corporation. Tutti i diritti sono riservati.

C:\Users\LEI00068>jupyter notebook
```

Python hands on

Notebook Python_basics

Example of Datasets

Chapter 2: Data Preprocessing

Motivation

- Raw data rarely comes in the form and shape that is necessary for the optimal performance of a learning algorithm.
- Real-world data is often dirty containing outliers, missing values, wrong data types, irrelevant features, or non-standardized data.
- Field of Data Preprocessing is really problem depend, these notes report only limited examples

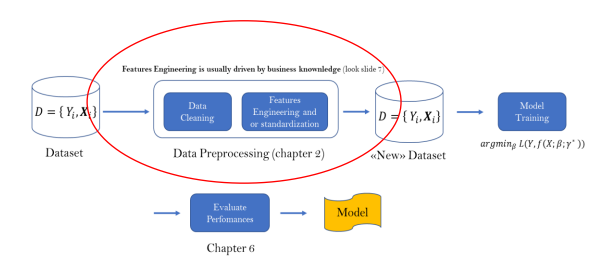
The success of a machine learning algorithm highly depends on the quality of the data fed into the model.



Transforming raw data into a useful format is an essential stage in the machine learning process.

Motivation

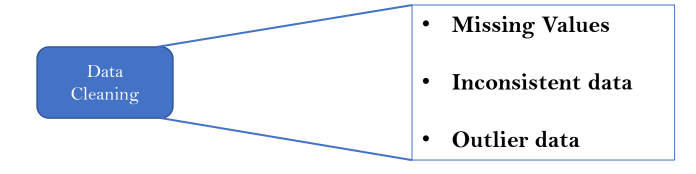
THIS IS YOUR MACHINE LEARNING SYSTEM? YUP! YOU POUR THE DATA INTO THIS BIG PILE OF LINEAR ALGEBRA, THEN COLLECT THE ANSWERS ON THE OTHER SIDE. WHAT IF THE ANSWERS ARE WRONG? JUST STIR THE PILE UNTIL THEY START LOOKING RIGHT.



We are here and will talk about:

- Data Cleaning
- Featuere Eng. and/or Standardization

Data Cleaning



The real-world data often has a lot of missing values due to failure in the recording process

Solution 1

Deleting the rows or columns having **null** values:

- If a columns have more than half of the rows as null then the column can be dropped.
- The rows which are having one value as null can also be dropped.
- Pandas can be used to easily handle null values (https://pandas.pydata.org/docs/reference/api/pandas.DataFrame.dropna.html)

Pros	Cons
Create a robust model	 If null are common in the dataset, a lot of information are removed
	• Removing a rows with a null cause the remove of good data

The real-world data often has a lot of missing values due to failure in the recording process

Solution 2

Imputing null values with statistics about that columns (e.g. mean/median)

• Pandas can be used to easily handle to impute null values (https://pandas.pydata.org/docs/reference/api/pandas.DataFrame.fillna.html)

Pros

No data loss, use all rows and columns

- Introduce synthetic data that may reduce model generalization
- Imputing method does not consider relation between features

The real-world data often has a lot of missing values due to failure in the recording process

Solution 3

Use model that automatically handle missing values e.g. XGBoost

Pros

- Null values are handled as "missing information"
- Only few model can handle null values

The real-world data often has a lot of missing values due to failure in the recording process

Solution 4

Encode null at particular value (e.g. -1 for positive features)

Pros

No data loss

Model may be not robust

Data Cleaning: Inconsistent data

The real-world data often has a lot of inconsistent values due to failure in the recording process:

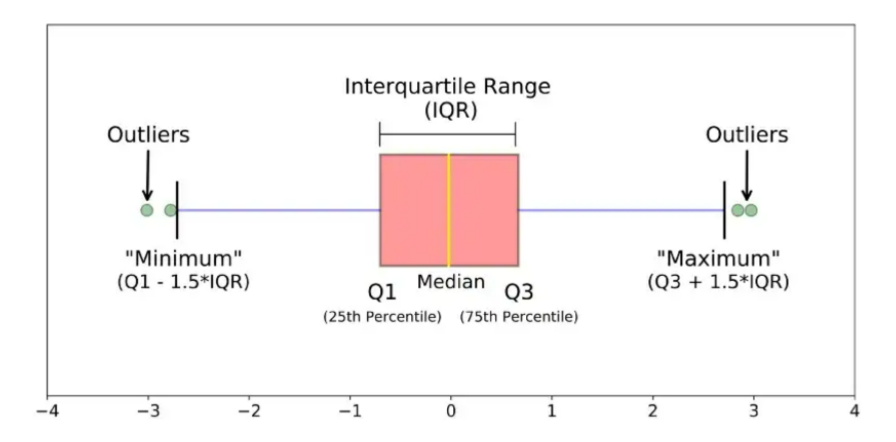
- Unfeasible data (too large or too low values ecc...)
- Data out of possible range (e.g. negative temperature in kelvin, negative ages, ecc...)

Solution

- Remove rows with inconsistent data
- **impute data** with suitable statistics
- use model that handle null values

Data Cleaning: Outlier

Data points that differ significantly from other observations. Some ML models may be sensitive to outlier, so they are excluded from the dataset



Data Cleaning: Outlier

Solutions 1

Remove outliers based on percentile rules, e.g. all data above 99 percentile.

Pros

• Create a model robust to outliers

- Some information are removed
- Univariate technique

Solutions 2

Use unsupervised model to detect outliers in the data (see chapter 7 or PCA at the end of this chapter)

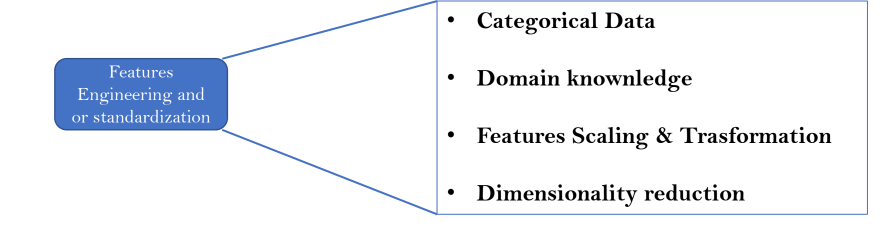
Pros

• Create a model robust to outliers

Model used to detect outlier may be limited

Feature Engineering

Feat. Eng. is the process of using domain knowledge to extract additional features from raw data



Feature Engineering: Categorical Data

Categorical data is a form of data that takes on values within a finite set of discrete classes

Some examples are:

- Province of birth or residence
- Gender
- Degree of education (phd, high school)

ML algorithms 'mainly'* work with numeric data \rightarrow convert/encode Categorical data into numerical data

Ordinal

maintain a natural order in their class of values



Ordinal Encoding

Non ordinal

do not maintain any natural/logical order



One-Hot Encoding

Feature Engineering: Categorical Data

Ordinal

maintain a natural order in their class of values

Ordinal Encoding

Higher class are mapped to higher class

- Phd ———•
- Master Degree 3
- Bachelor Degree • 2
 - High School • 1

Implemented in sklern: sklearn.preprocessing.OrdinalEncoder

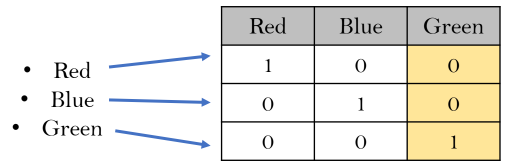
Non ordinal

do not maintain any natural/logical order

One-Hot Encoding

Create dummy variables.

For N class category \rightarrow give N binary (0/1) columns



Implemented in pandas & sklearn:

- sklearn.preprocessing.OneHotEncoder
 - Pandas.get_dummy

Feature Engineering: Categorical Data

What happens if the categorical data is the response y?



We are dealing with a classification task: $y \in \{0,1,...,m\}$



We can use LabelEncoder: sklearn.preprocessing.LabelEncoder

Encode target labels with value between 0 and $n_{class}-1\,$

For instance, a target for job status with 4 values:

- 0: Full-time
- 1: Part-time
- 3: Unemployed
 - 4: Intern

Feature Engineering: Domain knownledge

using domain knowledge to extract additional features from raw data

$$x_{new} = g(x_3, x_4, \dots)$$

Create a new predictor x_{new} as a function of other features

Improve the performances of ML model as compared to supplying only the raw data

The function g is obtained from domain knownledge

For instance, consider a dataset with x_1 : gain, x_2 : cost

$$x_3 = \frac{(gain - cost)}{cost}$$
 ROI: Return of Investement

Feature Engineering: Scaling

Some ML algorithms may need to scale features so that they are comparable:

- Compare **relative importance of features on a model**, e.g. in linear and logistic regression
- ML Methods involving the computing of a distance (eg. nearest neighborn, not studied in this course)
- Make features **independent from unit of measure** (usefull for PCA, next argouments)

Standardization

Average $\widehat{x_i} = \frac{(x_i - m_i)^2}{\sigma_i^2}$ Variance

Min-Max Scaling

Features max
$$\widehat{x_{-}i} = \frac{x_i - x_{max,i}}{x_{max,i} - x_{min,i}}$$
Features max

sklearn.preprocessing.StandardScaler

sklearn.preprocessing.minmax_scale

Feature Engineering: Dimensionality Reduction

Consider a dataset D with n rows (observations) on a set of p predictor, $x_1, ..., x_p$. When the values of p is large (e.g $p \gg 100$) it may be difficult to train a ML model from the fact that we need a **lot samples** n which, in principle, should **scale exponentially with** p (with p feature one may need $n = 10^p$ rows).

This problem is known as the **curse of dimensionality***



Need a model for **dimensionality reduction**: $p \to m$ with $m \ll p$

This technique should keep only the **relevant variables**

³⁵

Feature Engineering: Dimensionality Reduction

What are the **relevant variables**?

A lower-dimensional representation of *D* that contains as much as possible of the **variance** of the data: if we have a constant column (0 variance) it is not useful to make predictions

Principal Component Analysis (PCA) do exactly that!!

PCA aims to find a linear transformation from $x_1, ..., x_p$ to $z_1, ..., z_p$ such that:

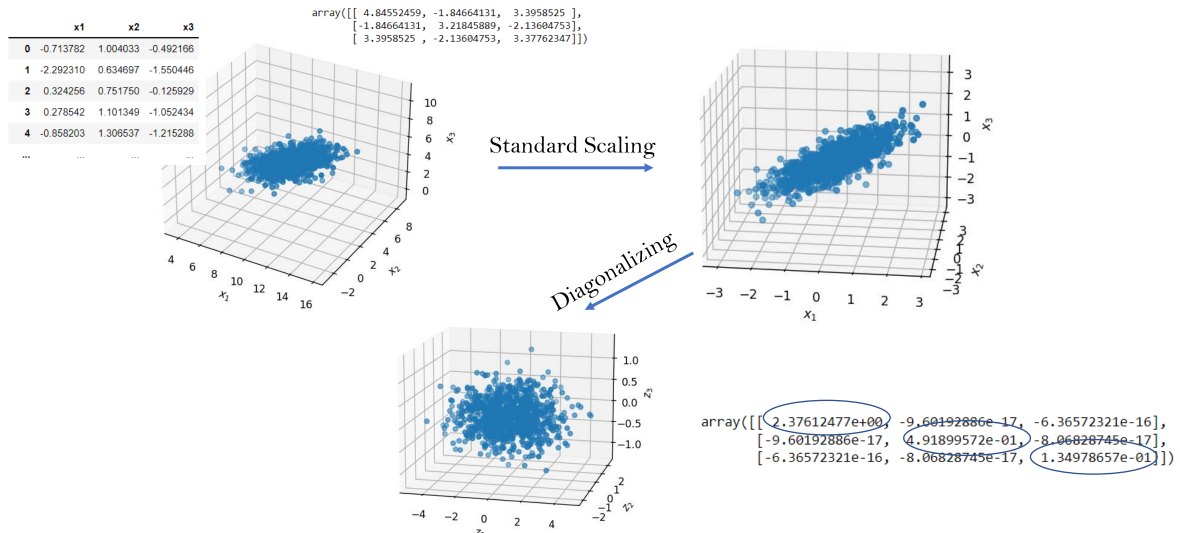
- The covariance matrix in the **z** space is diagonal (1)
- Keep only z that explain a given percentage of the total variance (2)

$$z_1 = \phi_{11}x_1 + \phi_{12}x_2 + \phi_{13}x_3$$
 $z_2 = \phi_{21}x_1 + \dots$
 $z_3 = z_3 = \dots$

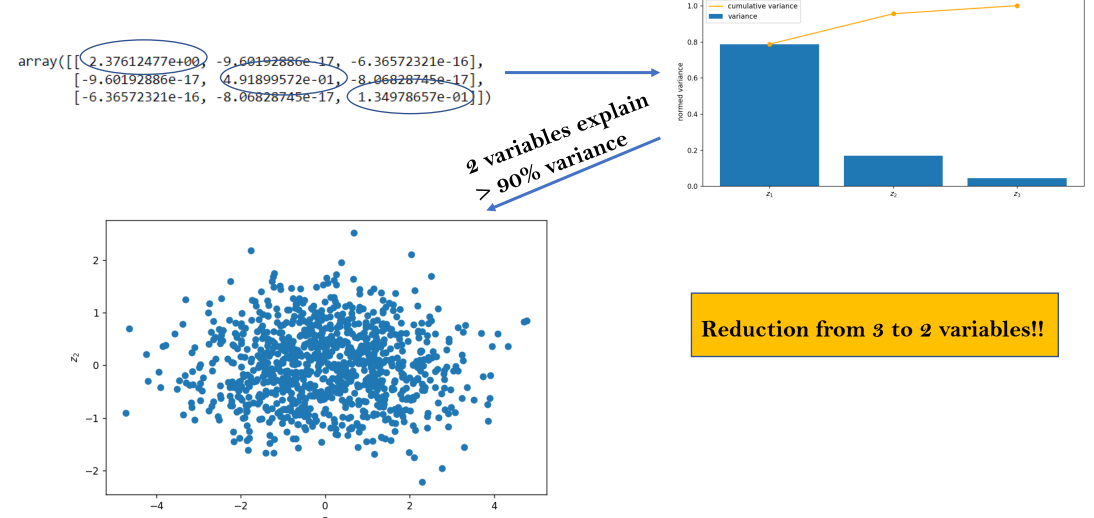
Find the matrix ϕ that solve (1) and then take (2)

3d example

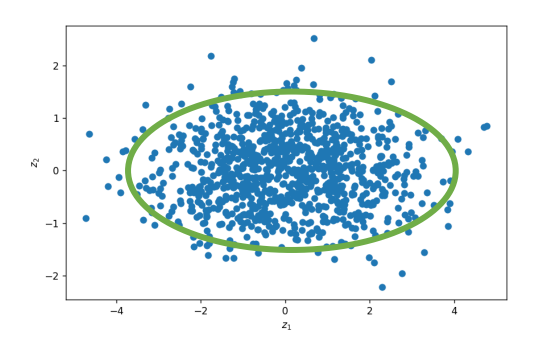
Feature Engineering: Dimensionality Reduction



Feature Engineering: Dimensionality Reduction



PCA for outliers detection



HP:

Outliers are point far from the center in the z-space

$$d = \sqrt{\sum_{i} \frac{z_i^2}{\sigma_i^2}}$$
 For each point

Mahalanobis distance: distance weighted with variance (covarince)

Note: in the z-space the covarince is diagonal otherwise formula is complex $d_M(\vec{x},Q) = \sqrt{(\vec{x}-\vec{\mu})^{\mathsf{T}}S^{-1}(\vec{x}-\vec{\mu})}$.

Define a threshold Z for d and label as outliers all points such that: d>Z

Choose Z based on some statistics about the distance compute on all the points, e.g. every points outside the green ellips is non an outlier

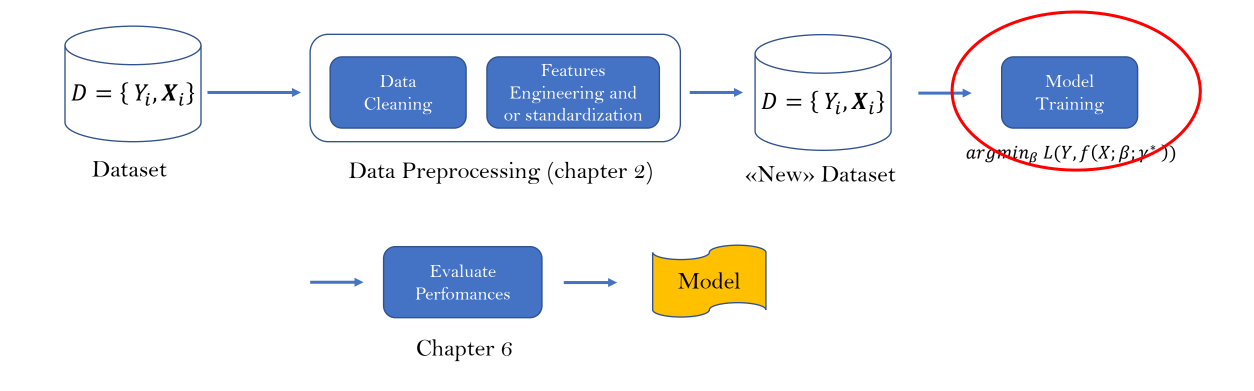
Python hands on

Notebook on cleaning data chapter 2

PCA example

Chapter 3: Linear Regression

Where we are



Linear Regression (LR)

The first (very simple) model we want to study is Linear Regression

In Linear Regression we suppose a linear relation between x_i and y

Consider a Dataset $D = \{x_i, y\}$ with p features and n rows (training dataset)

$$\hat{y} = f(x) = \beta_0 + \beta_1 x_1 + \dots + \beta_p x_p$$

 $\beta = [\beta_0, ..., \beta_n]$ are the parameter of the model LR have no hyper-parameter

$$\beta, \gamma = argmin_{\beta, \gamma} L(Y, f(X; \beta; \gamma))$$

Which L do we use? Mean Squared Error!!

$$L = \frac{1}{n} \sum_{i=0}^{n} (y_i - \hat{y}_i)^2 = \frac{1}{n} \sum_{i=0}^{n} (y_i - (\beta_0 + \beta_1 x_{i,1} + \dots + \beta_p x_{i,p}))^2$$
 y_i : the actual values \hat{y}_i : the predicted values

$$\beta^* = argmin_{\beta} \frac{1}{n} \sum_{i=0}^{n} \left(y_i - (\beta_0 + \beta_1 x_{i,1} + \dots + \beta_p x_{i,p}) \right)^2$$

Solve the optimization problem

$$\beta^* = argmin_{\beta} \frac{1}{n} \sum_{i=0}^{n} \left(y_i - \left(\beta_0 + \beta_1 x_{i+1} + \dots + \beta_p x_{i,p} \right) \right)^2$$

$$\beta^* = argmin_{\beta} \frac{1}{n} \left(Y - X\beta \right)^T (Y - X\beta)$$

$$\frac{d}{d\beta} L(\beta) = 0$$

$$\frac{d}{d\beta}(X\beta - Y)^T(X\beta - Y) = \frac{d}{d\beta}(\beta^T X^T X\beta - \beta^T X^T Y - Y^T X\beta + Y^T Y) =$$

$$= 2\beta^T X^T X - 2Y^T X = 2X^T X\beta - 2X^T Y$$

This is a scalar transpose has no effect \rightarrow we just prefer β at the end

Using Matrix Notation

$$\mathbf{y} = egin{bmatrix} y_1 \ y_2 \ dots \ y_n \end{bmatrix} \quad \mathbf{X} = egin{bmatrix} \mathbf{x}_1^\mathsf{T} \ \mathbf{x}_2^\mathsf{T} \ dots \ \mathbf{x}_n^\mathsf{T} \end{bmatrix} = egin{bmatrix} 1 & x_{11} & \cdots & x_{1p} \ 1 & x_{21} & \cdots & x_{2p} \ dots & dots & \ddots & dots \ 1 & x_{n1} & \cdots & x_{np} \end{bmatrix} \quad oldsymbol{eta} = egin{bmatrix} eta_0 \ eta_1 \ eta_2 \ dots \ eta_p \end{bmatrix}$$

The 1 are needed to fit the intercept eta_0

«Vector» derivatives

$$\alpha = \mathbf{y}^{\mathsf{T}} \mathbf{A} \mathbf{x} \qquad \frac{\partial \alpha}{\partial \mathbf{x}} = \mathbf{y}^{\mathsf{T}} \mathbf{A} \qquad \frac{\partial \alpha}{\partial \mathbf{y}} = \mathbf{x}^{\mathsf{T}} \mathbf{A}^{\mathsf{T}}$$
$$\alpha = \mathbf{x}^{\mathsf{T}} \mathbf{A} \mathbf{x} \qquad \frac{\partial \alpha}{\partial \mathbf{x}} = \mathbf{x}^{\mathsf{T}} \left(\mathbf{A} + \mathbf{A}^{\mathsf{T}} \right)$$

Different perspective: max log-likehood

Let us suppose that:

- y_i is a random variable distributed as a gaussian $G(\mu_i; \sigma)$
 - y_i are independent
- There is a linear relation between μ_i and x: $\mu_i(x) = \beta_0 + \beta_1 x_{i,1} + ... + \beta_p x_{i,p}$

The β can be computed using max likehood as follows:

The probability of observing the sequence $\{y_i\}$ $P(\{y_i\}) = \prod_{i=0}^n G(\mu_i; \sigma) = \prod_i^n e^{-\frac{(y_i - \mu_i(x))^2}{2\sigma^2}}$

Extraction the log
$$log P(\{y_i\}) = -\sum_{i=1}^{n} \frac{(y_i - \mu_i(x))^2}{2 \sigma^2}$$

 β are those maximazing the probability of observing the sequence (minimizing thus changing the sign)

$$\beta^* = argmin_{\beta} \sum_{i=0}^{n} (y_i - (\beta_0 + \beta_1 x_1 + ... + \beta_p x_p))^2$$

Same results, but different perspective!!

Evaluate a Linear Regression: Metrics

Mean squared erro	r
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$$MSE = \frac{1}{n} \sum_{t=1}^{n} e_t^2$$

Root mean squared error

$$RMSE = \sqrt{\frac{1}{n} \sum_{t=1}^{n} e_t^2}$$

Mean absolute error

$$MAE = \frac{1}{n} \sum_{t=1}^{n} |e_t|$$

Mean absolute percentage error

$$MAPE = \frac{100\%}{n} \sum_{t=1}^{n} \left| \frac{e_t}{y_t} \right|$$

Interpreting the coefficients

When the predictor are scaled the β can be used to understand the importance of the x_i over y

- Which is the most important feature? The x_i with the highest $|\beta_i|$ is the most important feature:
 - Positive β_i means that for positive values of x_i the y increase
 - Negative β_i means that for negative values of x_i the y increase
- The x_i with the lowest $|\beta_i|$ is the less important and (sometimes) can be removed from the dataset

Unluckly in real-world data this is often difficult to interpret variable importance. One problem is **collinearity** of predictors i.e., when one predictor can be expressed as a function (usually linear) of the other. In this case the estimated coefficients β may be wrong:

Actual relation $\hat{y} = 1 + 2x_1 + 3x_2$

Perfect collinearity
$$x_1 = 2x_2$$

$$\hat{y} = 1 + 7x_2$$

$$\hat{y} = 1 + \frac{7}{2}x_1$$

They all have the same performances of the actual model, but very different interpretation.

No more unique solution $(X^TX)^{-1}$ does not exist if there is collinearity :0 !!!

Ridge Regression

To avoid the collinearity and **improve performance and interpretability** of the model we need a **regolarizzation technique**

$$\hat{y} = f(x) = \beta_0 + \beta_1 x_1 + \dots + \beta_p x_p$$
 and should be fixed or computed in other ways (see chapter 6)
$$\beta^* = argmin_\beta \frac{1}{n} \sum_{i=0}^n \left(y_i - \left(\beta_0 + \beta_1 x_{i,1} + \dots + \beta_p x_{i,p} \right) \right)^2 + \lambda \sum_{j=0}^n \beta_j^2$$

This term try to keep the error between model and observation small

This term want to keep the β close to 0

 λ is a **hyperparameter** of the model

$$\beta^* = (X^T X + \lambda I)^{-1} X^T Y \rightarrow \hat{Y} = X_{new} \beta^* = X_{new} (X^T X + \lambda I)^{-1} X^T Y$$

This method select the relavant β_i (those different from 0) solving the collinearity problem: this can be also interpreted as a variable selection method!!

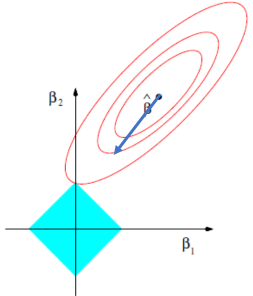
Other type of regolarizzation

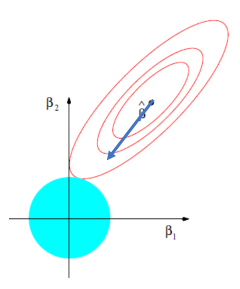
Lasso Regression

Ridge Regression

ElasticNet Regression

$$\beta^* = argmin_{\beta} \sum_{i=0}^{n} \left(y_i - (\beta_0 + \dots + \beta_p x_p) \right)^2 + \lambda \sum_{j=0}^{n} |\beta_j| \qquad \beta^* = argmin_{\beta} \sum_{i=0}^{n} \left(y_i - (\beta_0 + \dots + \beta_p x_p) \right)^2 + \lambda \sum_{j=0}^{n} \beta_j^2 \qquad \beta^* = argmin_{\beta} \sum_{i=0}^{n} \left(y_i - (\beta_0 + \dots + \beta_p x_p) \right)^2 + \lambda \sum_{j=0}^{n} \beta_j^2 \qquad \mathbf{Two\ hyperparameter}$$





If β^* is the solution of the non-regolarized problem, the regularization move the solutions toward the center of the axis (0): higher is λ more is close to 0

Polynomial Regression (PR)

What if the relation between x and y?

For the sake of simplicity, Consider a Dataset $D = \{x_1, y\}$ with 1 features and n rows and suppose:

$$\hat{y} = f(x) = \beta_0 + \beta_1 x_1 + \beta_2 x_1^2 + \beta_3 x_1^3 + \cdots$$

Polynomial Regression

x_1	y
1	12
4	17
3	45
	•••

Sklearn PolynomialFeatures

x_1	x_{1}^{2}	x_1^3	у
1	1	1	12
4	16	64	17
3	9	27	45

After the transformation, Poly Reg. is equivalent to Liner Reg. → Poly Reg. is just a feature eng. ☺

$$\beta^* = argmin_{\beta} \frac{1}{n} \sum_{i=0}^{n} \left(y_i - (\beta_0 + \beta_1 x_1 + \dots + \beta_p x_p) \right)^2 + \lambda \sum_{j=0}^{n} |\beta_j|$$

Kernel Regression

Interesting! we used a linear regression on a transformation of x to get a non-linear model with no effort! Let us try to generalize this approach:

$$x \in R^p \ \to \ \phi(x) \in R^m$$

Define a function ϕ that map an element in the p-dim features space to a m-dim space (like feature eng.)

Apply all formulas for Ridge* Linear Regression on $\phi(x)$

$$\hat{Y}(X_{new}) = X_{new}\beta = X_{new}(X^TX + \lambda I)^{-1}X^TY = Y^T(XX^T + \lambda I)^{-1}XX_{new}^T$$

Used identity abouy matrix

$$\hat{Y}(X_{new}) = Y^{T}(\phi\phi^{T} + \lambda I)^{-1}\phi\phi^{T}(X_{new})$$

 ϕ appears always in the form $\phi(X)\phi^T(X) \to \text{We can define a function K (kernel)}$ such as $K = \phi(X)\phi^T(X)$

In practice we do not need to explicit the function $\phi(x)$, but just K(x,y) (This is named the **kernel trick**)

Kernel Regression

$$\widehat{Y}(X_{new}) = Y^{T}(K + \lambda I)^{-1}k(x, X_{new})$$

$$K = \begin{pmatrix} \phi(x_1)\phi^T(x_1) & \cdots & \phi(x_n)\phi^T(x_1) \\ \vdots & \ddots & \vdots \\ \phi(x_1)\phi^T(x_n) & \cdots & \phi(x_n)\phi^T(x_n) \end{pmatrix} = \begin{pmatrix} K(x_1, x_1) & \cdots & K(x_1, x_n) \\ \vdots & \ddots & \vdots \\ K(x_n, x_1) & \cdots & K(x_n, x_n) \end{pmatrix}$$

Example 1 : Polynomial Kernel

$$x = (x_1, x_2) \longrightarrow \phi(x) = (1, \sqrt{2} x_1, \sqrt{2} x_2, \sqrt{2} x_1 x_2, x_1^2, x_2^2)$$
$$K(x, y) = (1 + xy^T)^2$$

Similar to 2-order polynomial regression, but with interaction between features

Example 2 : Gaussian Kernel

$$K(x,y) = e^{rac{|x-y|^2}{\sigma^2}} \qquad \qquad \phi_{RBF}(x) = e^{-\gamma x^2} igl[1, \sqrt{rac{2\gamma}{1!}} x, \sqrt{rac{(2\gamma)^2}{2!}} x^2, \sqrt{rac{(2\gamma)^3}{3!}} x^3, \ldots igr]^T,$$

Here ϕ is difficult to write and map x to infinite dimensional space !!!! (hint: series expansion)

Python hands on

Notebook on regression

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Chapter 4: Classification Methods

Logistic Regression (LR)

The first (very simple) classification model is Logistic Regression

Consider a Dataset $D = \{x_i, y\}$ with p features and n rows where $y \in \{0,1\}$

Instead of modelling the response, we model the probability p(x) that the record belong to a category

In Logistic Regression we have that:

$$p(y = 1 \mid x) = \frac{e^{(\beta_0 + \beta_1 x_1 + \dots + \beta_p x_p)}}{1 + e^{(\beta_0 + \beta_1 x_1 + \dots + \beta_p x_p)}}$$

$$p(y = 1 \mid x) = \frac{e^{(\beta_0 + \beta_1 x_1 + \dots + \beta_p x_p)}}{1 + e^{(\beta_0 + \beta_1 x_1 + \dots + \beta_p x_p)}}$$

$$= \frac{1}{1 + e^{(\beta_0 + \beta_1 x_1 + \dots + \beta_p x_p)}}$$

$$\log \frac{p(y = 1 \mid x)}{p(y = 1 \mid x)} = \beta_0 + \beta_1 x_1 + \dots + \beta_p x_p$$
Linear relation on logit

$$\log \frac{p(y-1|x)}{p(y=0|x)} = \beta_0 + \beta_1 x_1 + \dots + \beta_p x_p$$

Linear relation on logit!

Known as Logistic Function

Why this "strange" function and not a simple linear relation for p(x)?

Because in this way $p(x) \in [0,1]$ which is "natural" for probabilities. Many functions may meet this criteria, but here Logistic Function is used!

Logistic Regression (LR)

Now we need to computed the coefficients $\beta \rightarrow \text{Use Max. likehood}$

The probability of observing the sequence $\{y_i\}$ $P(\{y_i\}) = \prod_{y_i=1} p(y=1|x) \prod_{y_i=0} (1-p(y=1|x))$

$$\log P(\{y_i\}) = \sum_{i} y_i \ln p(y = 1|x) + (1 - y_i) \ln(1 - p(y = 1|x))$$

Substituting the form for p(y = 1|x) used in Logistic Regression, and using scalar product we obtain

$$\log P\left(\{y_i\}\right) = \sum_{i} y_i \beta \cdot x_i - \log(1 + e^{\beta \cdot x_i})$$

$$\beta^* = argmin_{\beta} - \sum_{i} y_i \beta \cdot x_i - \ln(1 + e^{\beta \cdot x_i})$$

Regolarizzation can be applyed

Changed sign to write as a min-problem

$$L(Y, f(X; \beta))$$
 Loss Function

Logistic Regression (LR)

$$\beta^* = argmin_{\beta}L(Y, f(X; \beta)) = argmin_{\beta}(-\sum_{i} y_i \beta \cdot x_i - \ln(1 + e^{\beta \cdot x_i}))$$
$$\frac{d}{d\beta_j}L(Y, f(X; \beta)) = 0 \rightarrow \sum_{i} -y_i x_j + x_j p(x_i; \beta) = 0$$

This is a **non-linear system** for all β_i and can **not** be solved analytically



Let's do it numerically: Gradient Descent!

Gradient Descent

We want to solve the problem: $\beta^* = argmin_{\beta}L(\beta)$ or equivalently, find β^* such as $\frac{d}{d\beta}L(\beta)|_{\beta^*} = 0$

Define an iterative (pseudo-dynamics) algorithm as follow:

$$\beta_{0} = \beta_{0}$$

$$\beta_{t+1} = \beta_{t} - \gamma \frac{d}{d\beta} L(\beta) \Big|_{\beta_{t}}$$

$$\downarrow \qquad \qquad \downarrow \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \downarrow \qquad \qquad \downarrow$$

 γ is the learning rate (lr) (delta-time in physics):

- The correct value depend on the problem at hand, i.e. it depends on the magnitude of $\frac{d}{d\beta}L(\beta)$
- The convergence of the algorithm critically depends on γ :
 - too small cause the process to get stuck
 - too large may cause instability

Gradient Descent: an example

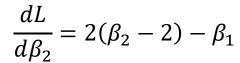
$$L = (\beta_1 - 1)^2 + (\beta_2 - 2)^2 - \beta_1 \beta_2$$

$$\frac{dL}{d\beta_1} = 2(\beta_1 - 1) - \beta_2$$

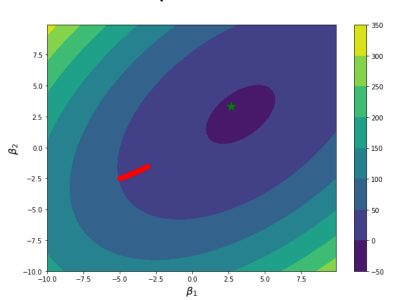
$$\frac{dL}{d\beta_1} = 2(\beta_1 - 1) - \beta_2$$

$$\beta_{t+1} = \beta_t - \gamma \frac{d}{d\beta} L(\beta) \Big|_{\beta_t}$$

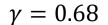
$$\gamma = 0.01$$

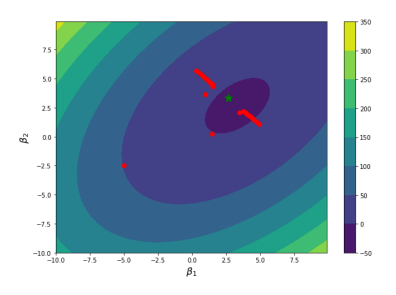


$$\gamma = 0.01$$



 γ is too small \rightarrow slow convergence





 γ is too big \rightarrow instability, bounce!

 $\gamma = 0.3$

5.0

2.5

-2.5

-5.0

-7.5

Interpreting the coefficients

When the predictor are scaled the β can be used to understand the importance of the x_i over y

- The x_i with the **highest** $|\beta_i|$ is the **most important**:
 - Positive β_i means that for positive values of x_i the βx increase, hence $p(x) \to 1$
 - Negative β_i means that for positive values of x_i the βx decrease, hence $p(x) \to 0$
- The x_i with the lowest $|\beta_i|$ is the less important and can be removed from the dataset
- Instead of p(x) you can interpret coefficients in term of odds i.e., $\ln\left(\frac{p}{1-p}\right)$

Like discussed for Linear Regression, also for logistic regression collinearity of predictors i.e., is a problem in the estimation of β coefficients. This problem can be solved as well applying reguralizzation techniques.

Logistic regression, given x_i output a probability p(x) that the record belong to the class 1. We could classify an observation based on some rules on p(x): let us define a threshold t such as:

- If p(x) > t the observation belong to class 1
- If $p(x) \le t$ the observation belong to class 0

One can be tempt to choose t = 0.5, but this value really depends on the problems at hand

Confusion Matrix

	Predicted Negative	Predicted Positive
Outcome Negative	True Negative(TN)	False Positive (FP)
Oucome Positive	False Negative(FN)	True Positive (TP)

$$Accuracy = \frac{TN + TP}{TN + FP + FN + TP}$$

Overall performance of the model: How many corrected classification over all observations

Logistic regression, given x_i output a probability p(x) that the record belong to the class 1. We could classify an observation based on some rules on p(x): let us define a threshold t such as:

- If p(x) > t the observation belong to class 1
- If $p(x) \le t$ the observation belong to class 0

One can be tempt to choose t = 0.5, but this value really depends on the problems at hand

Confusion Matrix

	Predicted Negative	Predicted Positive
Outcome Negative	True Negative(TN)	False Positive (FP)
Oucome Positive	False Negative(FN)	True Positive (TP)

$$recall = \frac{TP}{TP + FN}$$

Recall: How many good prediction over the total actual positive

Logistic regression, given x_i output a probability p(x) that the record belong to the class 1. We could classify an observation based on some rules on p(x): let us define a threshold t such as:

- If p(x) > t the observation belong to class 1
- If $p(x) \le t$ the observation belong to class 0

One can be tempt to choose t = 0.5, but this value really depends on the problems at hand

Confusion Matrix

	Predicted Negative	Predicted Positive
Outcome Negative	True Negative(TN)	False Positive (FP)
Oucome Positive	False Negative(FN)	True Positive (TP)

$$precision = \frac{TP}{TP + FP}$$

Precision: How many good prediction over the total predicted positive

Logistic regression, given x_i output a probability p(x) that the record belong to the class 1. We could classify an observation based on some rules on p(x): let us define a threshold t such as:

- If p(x) > t the observation belong to class 1
- If $p(x) \le t$ the observation belong to class 0

One can be tempt to choose t = 0.5, but this value really depends on the problems at hand

Confusion Matrix

	Predicted Negative	Predicted Positive
Outcome Negative	True Negative(TN)	False Positive (FP)
Oucome Positive	False Negative(FN)	True Positive (TP)

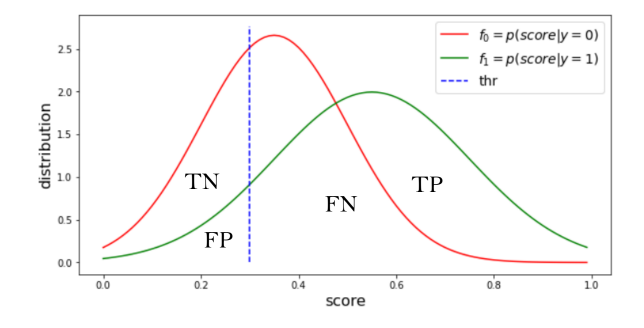
$$F_{beta} \ score = (1 + \beta^2) \frac{precision * recall}{\beta^2 precision + recall}$$

Precision: weight precision & recall F_1 score is harmonic mean of prec & recall

ROC

All the metrics discussed above depend on the choice of the threshold t. Here we want to find define a metric independent of t.

Actual Outcome	Score/Probability
О	0.2
0	0.8
О	0.15
0	0.25
1	0.12
1	0.81
1	0.6

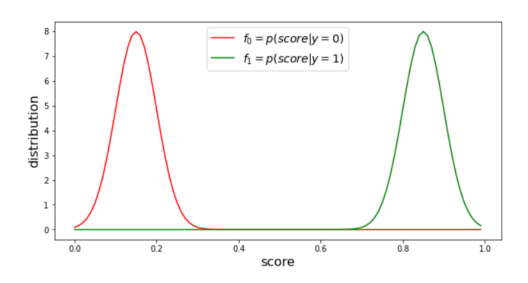


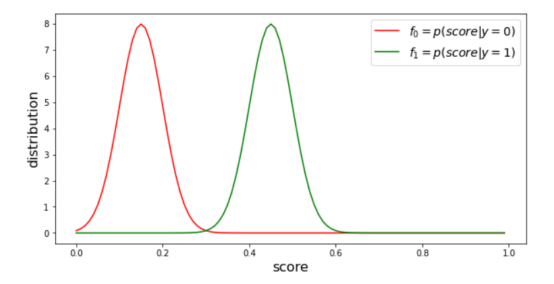
We can say that a good classifier output a score that can separate 0-class to 1-class

ROC

All the metrics discussed above depend on the choice of the threshold t. Here we want to find define a metric independent of t

Which can be a good metric?





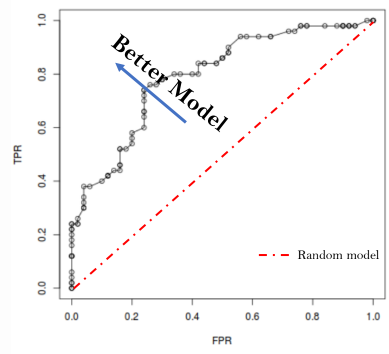
Both are very good classifier, only the scale of score/predictions change.

We need a metrics that measure how much this two distribution overlap

ROC

To define this metrics, firstly we define the ROC curve (Receiver Operating Characteristic). For each values of the threshold t, we compute: $TPR(t) = \frac{TP}{\#actual\ positive}$, $FPR(t) = \frac{FP}{\#actual\ negative}$

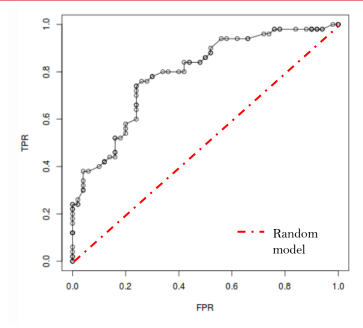
ROC curve is the plot of TRP (recall) against FPR

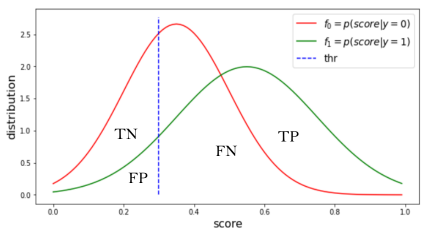


An example or ROC curve

- The point (0,1) is the perfect model!
- The red line is the random model line. Good model should have a ROC above this line (see proof in the upcoming slide)
- If a ROC is below the random model line, just inverting prediction 0 with 1 you obtain a better model above the red dashed line
- Given 2 models such as $ROC_1 > ROC_2$ for every FPR then model 1 is better than model 2

Area under the ROC

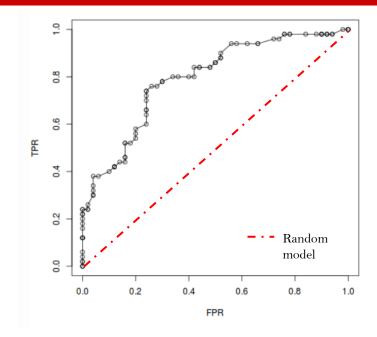


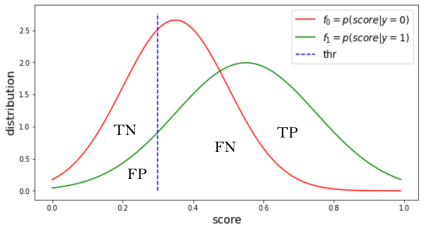


$$TPR(t) = P[p(x) > t \mid y = 1] = \int_{t}^{1} f1(t)dt$$
 $FPR(t) = P[p(x) < t \mid y = 0] = \int_{0}^{t} f_{0}(t)dt = F(t)$
 $p(x)$ is the score

$$\begin{aligned} &\text{AUC} = \int_0^1 T(F_0) \, \mathrm{d} F_0 & \text{Area Under the curve} \\ &= \int_0^1 P[\hat{p}(\mathbf{x}) > t \quad | y(\mathbf{x}) = 1] \, \mathrm{d} F_0 \\ &= \int_0^1 P[\hat{p}(\mathbf{x}) > t \quad | y(\mathbf{x}) = 1] \cdot \frac{\partial F(t)}{\partial t} \, \mathrm{d} t \\ &= \int_0^1 P[\hat{p}(\mathbf{x}) > t \, | y(\mathbf{x}) = 1] \cdot P[\hat{p}(\mathbf{x}') = t \, | y(\mathbf{x}') = 0] \, \mathrm{d} t \\ &= \int_0^1 P[\hat{p}(\mathbf{x}) > \hat{p}(\mathbf{x}') \, \& \, \hat{p}(\mathbf{x}') = t \, | y(\mathbf{x}) = 1 \, \& \, y(\mathbf{x}') = 0] \, \mathrm{d} t \\ &= P[\hat{p}(\mathbf{x}) > \hat{p}(\mathbf{x}') \, | \, y(\mathbf{x}) = 1 \, \& \, y(\mathbf{x}') = 0], \end{aligned}$$

Area under the ROC





$$TPR(t) = P[p(x) > t \mid y = 1] = \int_{t}^{1} f1(t)dt$$

$$FPR(t) = P[p(x) < t \mid y = 0] = \int_{0}^{t} f_{0}(t)dt = F(t)$$

$$AUC = \int TPR(FPR)dFPR = P[p(x) > p(x') | y(x) = 1 \& y(x') = 0]$$

In other word, given a randomly observation x belonging to class 1, and a randomly chosen observation x' belonging to class 0, the AUC is the probability that the classification algorithm will assign a higher score to x than to x':

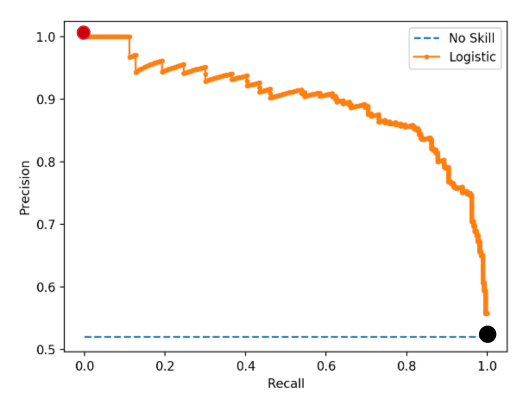
This is equivalent to ask that the red and green distribution are separated. Higher is the AUC, higher is the separation!

Precision Recall curve

Another widley used curve is the **Precion-Recall PR curve**.

For each values of the threshold t, we compute: precision(t), recall(t)

PR curve is the plot of precision vs recall



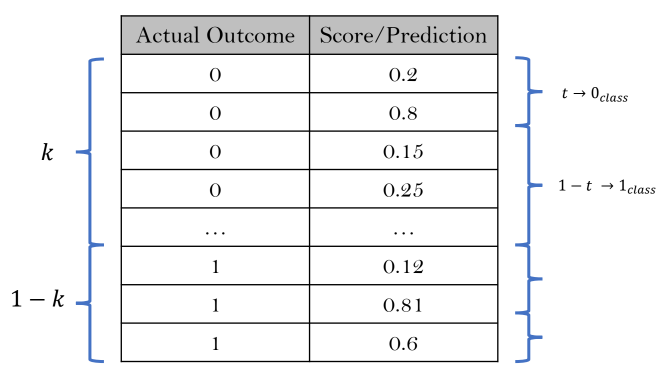
- The blue line is the random model line. Good model should have a PR above this line. Random model is the line $precision = \frac{\#total\ positive}{\#total\ case}$ fraction of positive case
- AUC PR is a valid metric, but has not easy probabilistic interpretation
 - **Dummy model** all predictes as class 1: recall = 1, precision = fraction of positive
 - **Dummy model** just one obs as class 1: recall = 1/N, precision = 100%

An example or ROC curve

ROC – PR: property of random model

The random model assign a score according to a uniform distribution between 0 and 1

N



 $N:observation, \quad k:fraction\ of\ 0-class \ t:threshold$

$$FPR = \frac{FP}{N} = \frac{Nk(1-t)}{Nk} = 1-t$$

$$TPR = \frac{TP}{P} = \frac{N(1-k)(1-t)}{N(1-k)} = 1-t$$

This is a straight line in *FPR* vs *TPR* plane

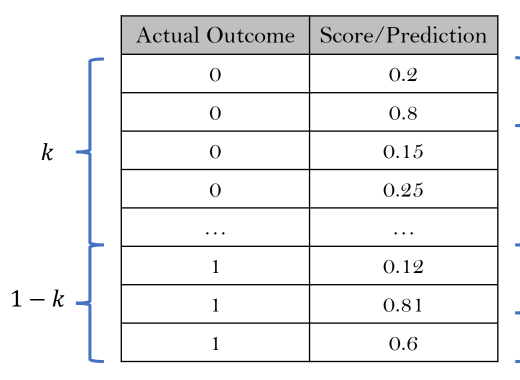
ROC – PR: property of random model

The random model assign a score according to a uniform distribution between 0 and 1

 $t \to 0_{class}$

 $1-t \rightarrow 1_{class}$

N



 $N:observation, \quad k:fraction\ of\ 0-class \ t:threshold$

$$precision = \frac{N(1-k)(1-t)}{Nk(1-t) + N(k-1)(1-t)} = (1-k)$$

(1-k) is the fraction of positive case and is independent on recall and t

Unbalanced Dataset

Unbalanced Dataset when the number of observation of 1 class is lower than those of 0 class (1% or 0.1%):

- Accuracy in not a good metrics to evaluate a model (see e.g. if the model predict all 0-class)
- Model prediction are often not calibrated, i.e. tends to be closer to 0
- AUC-ROC (but also tend to be too optimistic) and, mainly, precision-recall are the right way to measure performances (usually we are interested in 1-class related performaces)
- You can add weight on loss for 1 class to penalize more the 1 class term (there is an option in .fit() method)

$$\sum_{i} y_{i} \ln p(y = 1|x) + w_{i}(1 - y_{i}) \ln(1 - p(y = 1|x))$$

- Downsample 0-class observation
- Oversample 1-class observation (exists particular methods, e.g. SMOTE:out of the scope of this course)

More on precision & recall

Confusion Matrix

$$recall = \frac{TP}{TP + (FN)}; precision = \frac{TP}{TP + (FP)}$$

	Predicted Negative	Predicted Positive
Outcome Negative	True Negative(TN)	False Positive (FP)
Oucome Positive	False Negative(FN)	True Positive (TP)

Some example:

- For detect a bomb at an airport, anything that doesn't account for **false-negatives** is a problem. **Recall** is a better measure than precision. Same for medical application (e.g., cancer modelling)
- For YouTube recommendations, false-negatives is less of a concern. Precision is better here.
- If I am predicting if a patient should be given a medicine, that when given to healthy person is catastrophic: we need to avoid predicting a positive for a negative instance. Since we want to avoid FP errors, we must prefer **Precision**
- Usually, you can assign a business value to FP and FN (thus to prediction and recall) and just compute this metrics.

Multinomial logistic regression

Consider a Dataset $D = \{x_i, y\}$ with p features and n rows where $y \in \{1, ..., n\}$ y can be interpreted as the result of label encoder, i.e. categories

$$p(y = 1 \mid x) = \frac{e^{\beta_1 x}}{\sum_k e^{\beta_k x}}$$

$$p(y = 2 \mid x) = \frac{e^{\beta_2 x}}{\sum_k e^{\beta_k x}}$$

•••

$$p(y = k | x) = 1 - \sum_{i} p(y = i | x)$$

One β vector for each class usually represented as a matrix

Softmax Function

$$\log P(\{y_i\}) = \sum_{i} \sum_{k} I(y_i = k) \log p(y_i = k \mid x)$$

Generalize Loss for binary variables



Minimized by gradient descent

Generalized Linear Model

Distribution	Target Domain	Model hp	Likehood	Loss
Bernoulli	<i>y</i> ∈ {0,1}	$p(x) = \frac{1}{1 + e^{\beta x}}; \log \frac{p}{1 - p} = \beta x$	$P(\{y_i\}) = \prod_{y_i=1} p \prod_{y_i=0} (1-p)$	$L = \sum_{i} y_{i} \beta x_{i} - \ln(1 + e^{\beta x_{i}})$
Poisson / Count	$y \in \{0,1, \dots n\}$	$\ln(\mu(x)) = \ln(E(y x)) = \beta x$	$P(\{y_i\}) = \prod_i \frac{\mu^{y_i}}{y_i!} e^{-\mu}$	$L = \sum_{i} y_{i} \beta x_{i} - e^{\beta x_{i}}$

This approach can be generalized to different distribution:

- Suppose a probability distribution for y_i
- Define a relation between x and some moments of y_i
- Compute the probability of the sequences, extract the log and compute the Loss

Python hands on

Notebook on regression

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Chapter 5: Evaluate a Model

Motivation

- In all the real-world situation, we do not really care how the model works well on the **training data**. Rather, we are interested in the performance of the model to previously **unseen test data**:
 - Performance on unseen test data are referred as the **generalization performance** of the model.
 - A model that perform well on training, but has low performance on test suffers from overfitting
 - Trade-off between generalization & overfitting is often referred as **bias-variance trade-off**.
- Moreover, we need also a procedure to compute the hyper-parameters of a given model (e.g., the value of λ in LASSO/RIDGE regression, or the kernel for a regression)

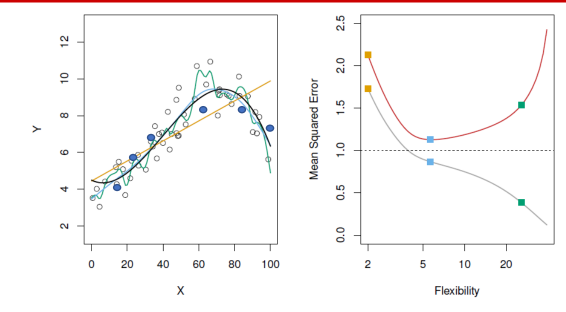


In this chapter we will introduce statistical methods aimed at evaluate overfitting and compute model hyper-parameter

Overfitting

Just consider two sets of data:

- Train data: used to build the model
- Test data: used to evaluate the model (must remain unseen)



Right) From linear regression, to high order polynomial regression. Training data are the empty dots.

Left) Performance on train data (gray line) improves as the model complexity increase while, performance on test data (red line) becomes lower. This is a fundamental property of statistical learning holding regardless of the datasets and of the model being used. Model with order equal to 20 overfits.

Idea: Seems reasonable to choose the model with polynomial order equal to 5: it show good performance for both sets (but we are selecting the model looking at data that should be unseen: data leak). We are discussing together the generalization performance and the choose of the hyper-parameter of the model. 80

Bias – Variance trade-off

It can be shown that the expected error on a new unseen points x_o can be written as a sum of 2 terms:

$$E\left(y_0 - \hat{f}(x_0)\right)^2 = \operatorname{Var}(\hat{f}(x_0)) + \left[\operatorname{Bias}(\hat{f}(x_0))\right]^2 \qquad \text{Mean squared error case}$$

The overall expected test MSE can be computed by averaging over all possible values of x_0 in the test set

Variance refers to the amount by which the estimated model f would change if we use a different training data set. In general, more **complex/flexible** statistical methods have higher variance.

Bias refers to the error that is introduced by approximating a real-world problem, by a much simpler model.

- in order to minimize the expected test error, we need to select a statistical learning method that achieves low variance and low bias but...
- For more **complex/flexible** methods, the variance will increase, and the bias will decrease. **Should find a trade-off**.

The simpler setup: train-test-validation splitting

If we are in a **data-rich situation**, the best approach is to randomly divide the dataset into three parts:

Training set	Validation set	Test set
Used to fit the model (70% of data)	Used to evaluate the model performances (15% of data)	Used to evaluate the model generalization (15% of data)

In practice: define a grid for all the hyper-parameter e.g., $\lambda \in \{\lambda_1, ..., \lambda_L\}$:

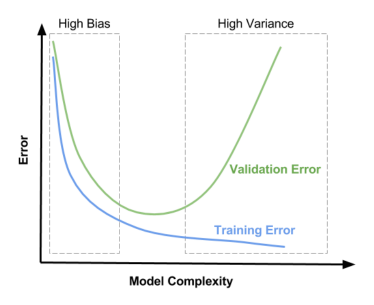
- For each value of λ , train the model on the train set
- Evaluate the performance of the model on the validation-set
- Choose the hyper-parameter λ^* with the **best** validation performance (why not optimization*??)
- Retrain the model with λ^* and evaluate the generalizzation error on the test. Finally compare performances between sets to adress possible overfitting!

This setup work well in a data-rich situation

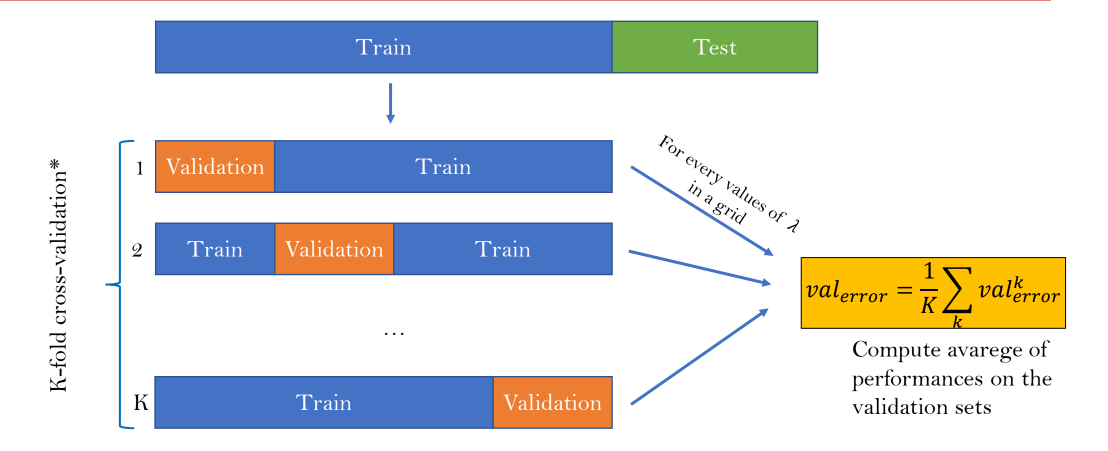
The simpler setup: train-test-validation splitting

If we are in a **data-rich situation**, the best approach is to randomly divide the dataset into three parts:

Training set	Validation set	Test set
Used to fit the model (70% of data)	Used to evaluate the model performances (15% of data)	Used to evaluate the model generalization (15% of data)



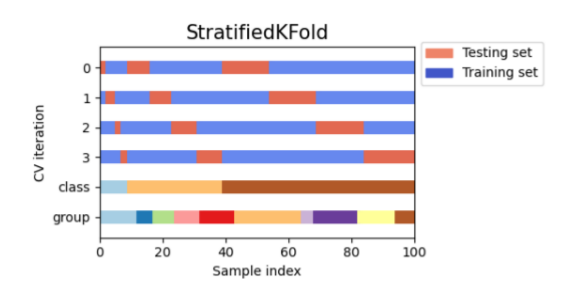
Cross-validation: less data-rich situation

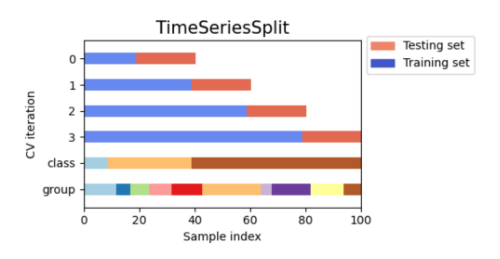


Choose the hyperparameter λ^* with the **best** cross-validation error! This approach is computationally expensive, but does not *waste* data (as the case for arbitrary validation)

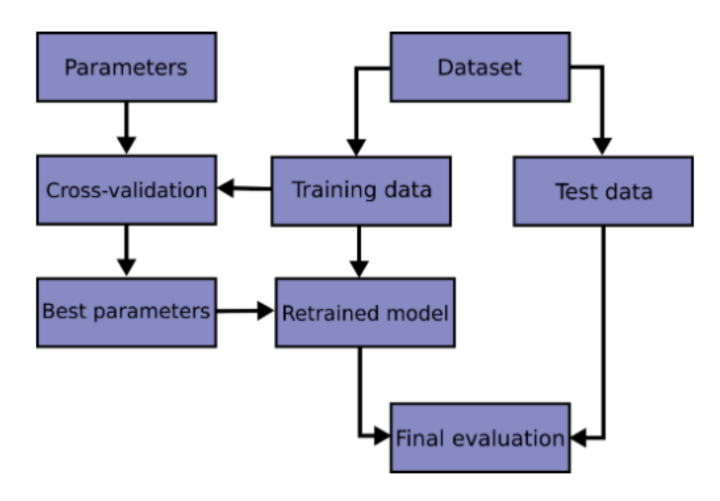
Cross-validation: special cases

- When applying cross-validation on a classification problem one may want that all the dataset contains approximately the same percentage of samples of each target class as the complete set: this can be done using **Stratified k-fold**
- In this course we do not study **time series model**, but here we want to highlight that in some case cross-validation can be tricky, like in the time-series model setup.





The general approach



Conclusions

- There can be some setup in which cross-validation may not work well, overestimating validation error performance.
- There exists other statistical tool (boostrapping methods, montecarlo sampling) which aim to better estimate validation error (for instance by better estimating its variance), but these methods are below the scope of this course.
- Scaling and any type of preprocessing should be fitted only on train dataset and the used on test data

Chapter 6: Tree-based Model

Motivation

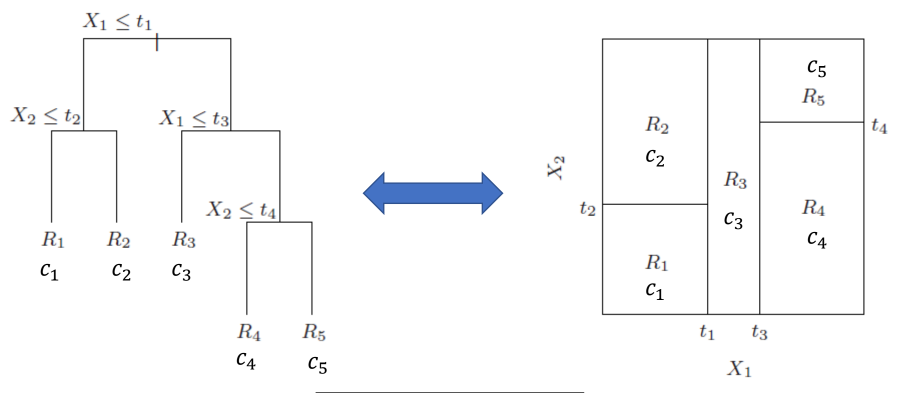
This chapter deals with a new class of model called tree-based model that are widley used both for regression and classification. We will study:

- Simple CART (Classification And Regression Tree)
- Random Forest
- Boosting tree
- Extreme Boosting Tree (XGBoost)

Finally we will introduce SHAP a tree-explainar model to understand the impact of the features over the final prediction.

CART - Regression

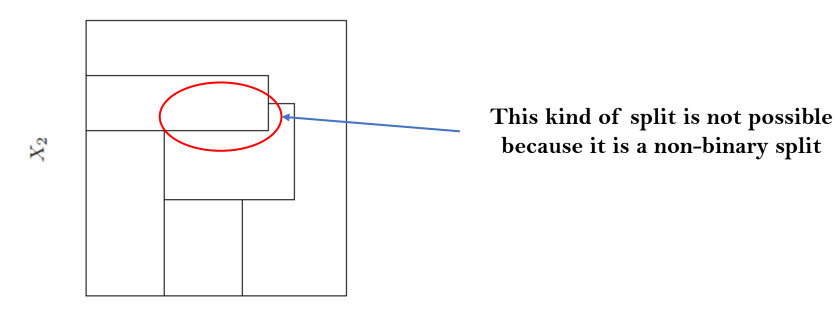
Tree-based methods partition the feature space into a set of rectangles (binary partitions), and then fit a simple model (like a constant) in each one



$$f(x) = \sum_{m=1}^{M} c_m I(x \in R_m)$$

CART - Regression

Tree-based methods partition the feature space into a set of rectangles (binary partitions), and then fit a simple model (like a constant) in each one



 X_1

CART - Regression

Consider a Dataset $D = \{x_i, y\}$ with p features and n rows (training dataset)

 $\beta, \gamma = argmin_{\beta, \gamma} L(Y, f(X; \beta; \gamma))$

$$f(x) = \sum_{m=1}^{M} c_m I(x \in R_m)$$

$$f(x) = \sum_{m=1}^{M} c_m I(x \in R_m) \qquad R_m^*, c_m^* = argmin_{R_m, c_m} \sum_{i=0}^{n} (y_i - f(x))^2$$
 (1)

For regression Mean Squared Error is a good Loss

Suppose first that we already have a partition into M regions $R_1^*, ... R_M^*$ hence:

$$\frac{d}{dc_m} \sum_{i=0}^n (y_i - f(x))^2 = \sum_{i \in R_m^*} 2(y_i - c_m) = 0 \to \sum_{i \in R_m^*} y_i - \#m \ c_m = 0$$

The coeff c_m compare only In the sum for the observation $i \in R_m^*$

$$c_m^* = \frac{1}{\#m} \sum_{i \in R_m^*} y_i = ave(y_i | x \in R_m^*)$$

The optimal region R_1^* , ... R_M^* are computed following a greedy optimization* algorithm that aim to solve (1)

CART – the greedy algorithm

The Region R is defined by a set of two informations: the feature and the split point $\{j, s\}$

Residual sum of square

Starting with all the data consider a feature j and a split point s and compute:

$$RSS = \sum_{i \in R_1} (y_i - c_1)^2 + \sum_{i \in R_2} (y_i - c_2)^2 \quad where \quad R_1 = \{x \mid x_j < s\} \text{ and } R_2 = \{x \mid x_j \ge s\} \text{ and } c_1 = ave(y_i \mid x \in R_1)$$

Cycling on all possible valus of (j, s) find those minimizing $RSS \rightarrow (j^*, s^*)$

Then one can repeat recursively the splitting process on each of the two regions.

How large should we grow the tree?

Regolarization and/or Cross Validation

CART – Regolarization

The process described above is likely to overfit the data, how to prevent this problem?

• **Regolarization**: Pruning the tree. This is tipically done by adding to the loss a term $+\lambda |T|$ where |T| is the number of leafs of the tree (terminal nodes) and find the best value of λ via cross-validation:

$$argmin_{R_m,c_m} \sum_{i=0}^{n} (y_i - f(x))^2 + \lambda |T|$$

• Cross-validation: the tree-depth is an hyperparameter of the model, it can be chosen via cross-validation

CART – Hyperparameter

The process described above is likely to overfit the data, how to prevent this problem?

- max_depth: The maximum depth of the tree (most important)
- min_samples_split: The minimum number of samples required to split an internal node
- min_samples_leaf: The minimum number of samples required to be at a leaf node

They can be chosen via cross-validation

Example: For instance, if min_samples_split = 5, and there are 7 samples at an internal node, then the split is allowed. But if the split results in two leaves, one with 1 sample, and another with 6 samples. If min_samples_leaf = 2, then the split won't be allowed (even if the internal node has 7 samples) because one of the leaves resulted will have less then the minimum number of samples required to be at a leaf node.

CART – Classification

A classification tree is very similar to a regression one. For a classification tree, the leaf prediction is the **most** commonly occurring class of training observations in the leaf-region to which it belongs, usually we are also interested in the class proportions among the training observations that fall into that region.

Use same growing algorithm as regression tree, but instead of RSS it consider a different split metrics:

Entropy (also called cross-entropy):

Gini Index:

 $Entropy = -p_1 \log p_1 - (1 - p_1) \log(1 - p_1)$ or: $Gini = p_1(1 - p_1)$

where p_1 is the 1-class proportion on the leafs

In essence, one have to compare the value of *Entropy* or Gini between parent and child leaf: if such difference is positive, i.e. E decrease, we can split.

CART – Classification

Entropy (also called cross-entropy):

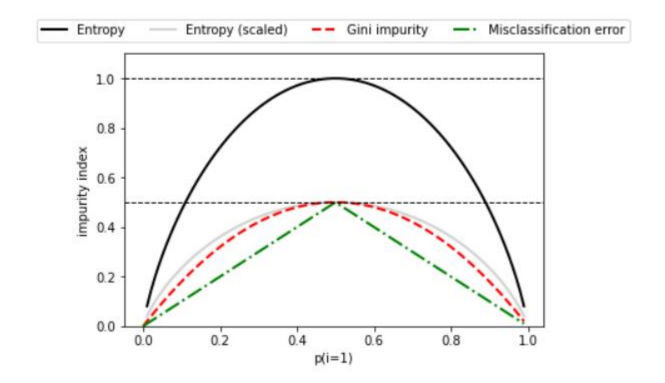
$$Entropy = -p_1 \log p_1 - (1 - p_1) \log(1 - p_1)$$

Gini Index:

$$Gini = p_1(1 - p_1)$$

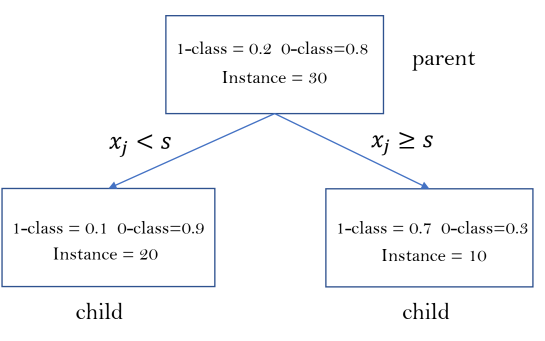
where p_1 is the 1-class proportion on the leafs

or:



CART – example

Introduce gain which measure how much the entropy (or gini) increase or decrease after a split



$$Gain = E_{parent} - E_{childs}$$

$$E_{parent} = -0.2 \log 0.2 - 0.8 \log 0.8 = 0.50$$

$$E_{childs} = \frac{20}{30} (0.1 \log 0.1 + 0.9 \log 0.9) + \frac{10}{30} (0.7 \log 0.70 + 0.3 \log 0.3) = 0.42$$

$$gain = E_{parent} - E_{child} = 0.08$$
 Gain is positive thus spit!

If **entropy decrese (gain is positive)** as the tree growth means that the uncertain about the class attribution decrease!

CART conclusions

Pros

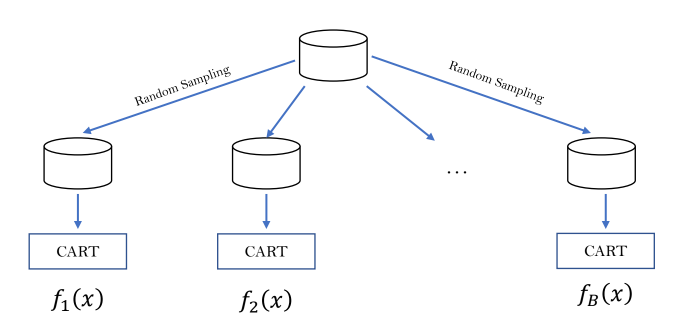
- Trees are very easy to explain
- Some researchers believe that decision trees more closely mirror human decision-making
- Trees can be displayed graphically and are easily interpreted even by a non-expert (especially if they are small).
- The feature to put at the root node is the one with the most influence on the outcome. This is essential for business interpretation of the results

- CART generally do not have the same level of predictive performances as some of the other regression and classification approaches seen in this course.
- CART can be non-robust: a small change in the data can cause a large change in the final estimated tree.

Random Forest (RF)

"Why build one tree when you can make a forest?"

The idea is to average many CART model, in order to improve performance



In order to decorrelate the tree and avoid very similar CART at each split in the tree the algorithm is allowed to consider a random minority of the available predictors! Indeed, averaging many highly correlated model does not lead to performance improvement and reduction in variance

$$f(x) = \frac{1}{B} \sum_{b=1}^{B} f_b(x)$$

For classification, majority voting is used.

RF – Hyperparameter

- n_estimators: the number of tree in the forest
- max_features: the number of features to consider during a split
- max_samples: the number of sample for random sampling
- max_depth: The maximum depth of the tree
- min_samples_split: The minimum number of samples required to split an internal node
- min_samples_leaf: The minimum number of samples required to be at a leaf node
- ccp_alpha: Complexity parameter used for Minimal Cost-Complexity Pruning.

Boosting tree

Boosting: build a model that combines the outputs of many weak model to have better result.

$$f(x) = \sum_{m} h_m(x; \gamma_m)$$

Where h_m are a very **simple** function: here a **very shallow** tree is used

As usual we aim to find a solution γ_m such as:

$$\gamma_m^* = argmin_{\gamma_m} \sum_i L(y_i, f(x))$$

The idea is to solve this problem in a gradient descent fashion

Boosting tree: gradient descent

$$f(x) = \sum_{m} h_m(x; \gamma_m) \to f_{m+1} = f_m + h_m$$
Iterative update like gradient descent

$$\beta_{t+1} = \beta_t - \gamma \frac{d}{d\beta} L(\beta) \Big|_{\beta_t}$$

$$h_m \to -\frac{d}{df_m} L(y, f_m) \Big|_{f_m}$$

Let us just build the model iteratively:

- f_0 is a constant ginven by $f_0 = argmin_{f_0} \sum_{i=0}^n (y_i f_0)^2 \rightarrow f_0 = ave(y_i)$
- For each observation compute $r_1 = -\frac{d}{df_0}L(y, f_0)|_{f_0}$
- Fit a regression tree $h_1(x; \gamma_1)$ on r_1
- Update $f_1(x) = f_0(x) + \epsilon f_1(x)$ where ϵ is an input parameter
- Iterate until $m \rightarrow M$

$$L = (y_i - f(x))^2$$

$$\frac{d}{df_0} L(y, f_0)|_{f_0} = \frac{d}{df_0} (y_i - f_0)^2$$

$$(y_i - f_0) \to r_1$$

residual of the previous step!!

Thus $h_1(x; \gamma_1)$ is fitted on the error made by previous predictor

Boosting tree – Hyperparameter

- n_estimators: the number of tree in the forest
- learning_rate ϵ : shrinks the contribution of each tree by learning_rate: $f(x) = f_0 + \epsilon f_1(x) + \dots$ There is a trade-off between learning_rate and n_estimators
- **subsample**: The fraction of samples to be used for fitting the individual base learners
- max_depth: The maximum depth of the tree
- min_samples_split: The minimum number of samples required to split an internal node
- min_samples_leaf: The minimum number of samples required to be at a leaf node
- ccp_alpha: Complexity parameter used for Minimal Cost-Complexity Pruning.

They can be chosen via cross-validation

Variable importance

Like in any other ML model, for tree-based model is important to understand the impact of the features on the labels. CART are highly interpretable: the model can be represented by a simple two-dimensional graph. Linear combinations of trees lose this important feature and must therefore be interpreted in a different way.

RF and Boosting critically improves prediction accuracy at the expense of interpretability.

We can measure how each feature decrease the *impurity* of the split (the feature with highest decrease is selected for internal node). For each feature we can collect how on average it decreases the Loss, i.e. the splitting criterion. The average over all trees in the model is the measure of the feature importance.

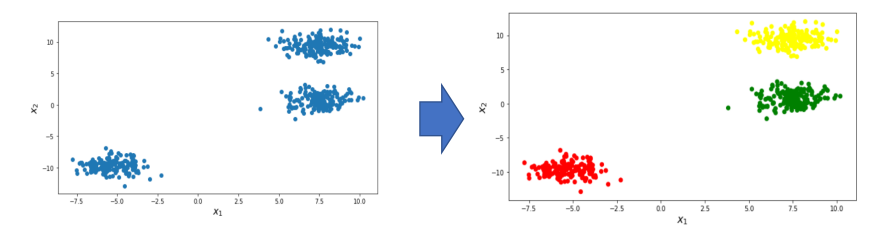
$$I_V(N) = \frac{1}{|S|^2} \sum_{i \in S} \sum_{j \in S} \frac{1}{2} (y_i - y_j)^2 - \left(\frac{|S_t|^2}{|S|^2} \frac{1}{|S_t|^2} \sum_{i \in S_t} \sum_{j \in S_t} \frac{1}{2} (y_i - y_j)^2 + \frac{|S_f|^2}{|S|^2} \frac{1}{|S_f|^2} \sum_{i \in S_f} \sum_{j \in S_f} \frac{1}{2} (y_i - y_j)^2 \right)$$

Another very popular library for variable importance is SHAP, but it is outside the scope of this course

Chapter 7: Unsupervised Method: Clustering

Unsupervised Learning

- Unsupervised learning algorithms work with data that isn't labelled, i.e no y is available
- It aims to partition data into distinct groups: observations within each group are "similar", while observations in different groups are "different": this is the definition of clustering. The heart of Clustering methods is to define what similar and different mean!
- **Example**: Clustering Customers in an online shop:
 - You run a large online store and you would like to personalize the user's shopping experience
 - Each user is described by a list of features
 - Clustering will divide your user base in different group: within each group users will have similar shopping pattern, so they may have similar ads.



K-means

Consider a Dataset $D = \{x_i\}$ with p features and n rows In the k-means alghorithm "similarity" is interpreted as "distance". The simplest distance measure is the Euclidean distance, using vector notation:

$$d(x,y) = (x - y)(x - y)^T$$

The output of k-means is a vector c_i of cluster assignment for each sample and k mean vectors μ_k such that:

$$c_i^*, \mu_k^* = argmin_{c,\mu} \sum_i \sum_k 1(c_i = k)(x_i - \mu_k)(x_i - \mu_k)^T$$

- The value of k is fixed (we will see later how to choose it)
- μ_k are the centroid of the cluster i.e., the "average" element that represent such cluster
 - Aim is to minimize the total distance between points and centroid
 - This suffer for the curse of dimensionality

K-means

$$c_i^*, \mu_k^* = argmin_{c,\mu} \sum_i \sum_k 1(c_i = k)(x_i - \mu_k)(x_i - \mu_k)^T$$

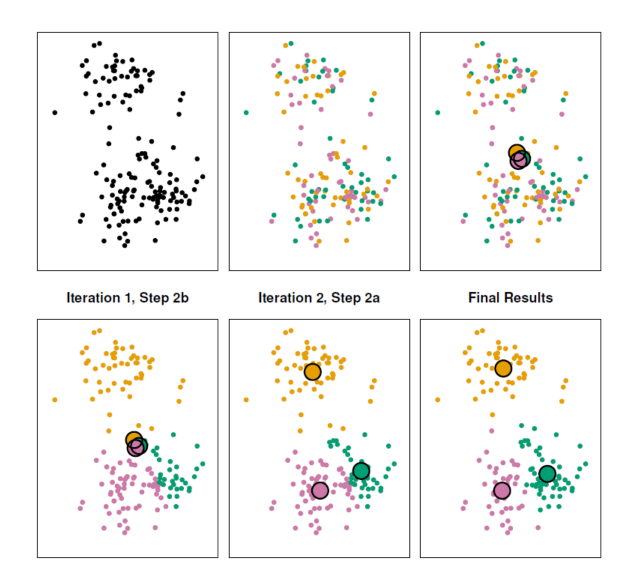
While this problem can be solved via gradient-descent, here we study a different algorithm that converge to the solutions of the problem:

- 1. Initialization: Randomly assign a cluster, from 1 to K, to each of the observations.
- 2. For each cluster, compute the cluster centroid as (center of mass, average):

$$\mu_k = \frac{1}{\#l} \sum_{l \in k} x_l$$

- 3. Assign each observation to the cluster whose centroid is closest, thus computing c_i
- 4. Repeat 2), 3) until c_i does not change. Output μ_k , c_i .

K-means

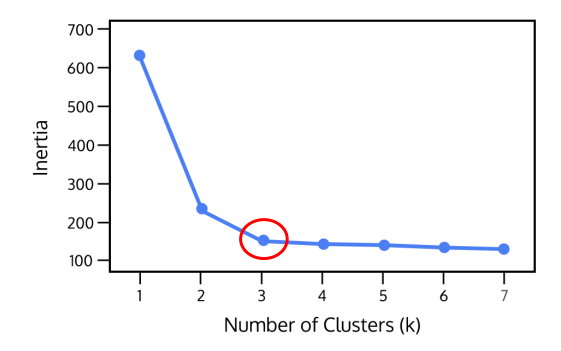


K-means

How to choose the correct number of cluster?

$$Inertia = \sum_{i} d_i^2$$

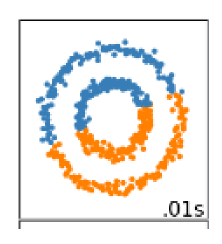
Compute the total distance to the cluster centroid for each value of k > This is the same as the optimal Loss Value



k = 3 is the optimal value since the Inertia does not decrease anymore

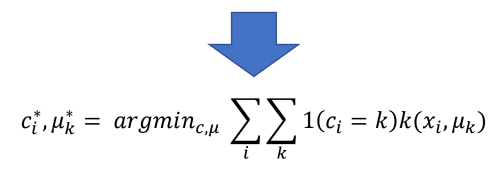
Inertia is also a performance metric

Kernelised K-means



With some dataset k-means could fail to extract the "correct" cluster. This is due to the definition of distance that has been used (for instance Euclidean).

This can be solved used a kernelized k-means algorithm.



Kernelised K-means

$$c_{i}^{*}, \mu_{k}^{*} = argmin_{c,\mu} \sum_{i} \sum_{k} 1(c_{i} = k)k(x_{i}, \mu_{k})$$

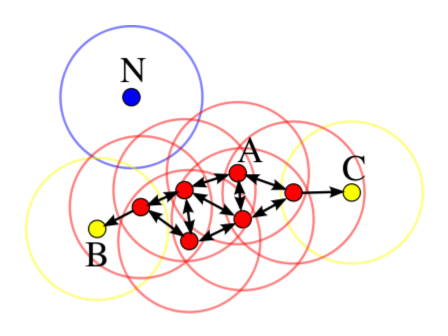
- 1. Initialization: Randomly assign a cluster, from 1 to K, to each of the observations.
- 2. For each cluster, compute the cluster centroid as (center of mass, average):

$$\mu_k = \frac{1}{\#l} \sum_{l \in k} \phi(x_l)$$

- 3. Assign each observation to the cluster whose centroid is closest, thus computing c_i
- 4. Repeat 2), 3) until c_i does not change. Output μ_k , c_i .

For this algorithm we need to access to the feature $\phi(x_l)$

DBSCAN



Density-based spatial clustering (DBSCAN) is a very popular clustering alghoritm

Some key idea are (defined two parameter MinPts and ϵ):

- red points are core points, because circle of ε -radius contain at least MinPts points (including the point itself).
- Yellow points are frontier points, and they belong to the cluster
- Point N is a noise point that is neither a core point nor directly-reachable.

- 1.DBSCAN does not require one to specify the number of clusters as opposed to k-means
- 2.DBSCAN can find arbitrarily-shaped clusters like those on previous slide
- 3.DBSCAN can be used as an outliers-detection methods:N-points are the outliers

Backup

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More on tree

Extreme Boosting tree

Same principles as Boostin method \rightarrow write f(x) as a sum of "weak" learner (shallow tree)

$$f(x) = \sum_{m} h_m(x; \gamma_m) \qquad f(x) = \sum_{m} h_m(x; \gamma_m) \to f_{m+1} = f_m + h_m$$

At the *T* stage we have:

$$L_T = \sum_{i} L(y_i, f_T(x)) = \sum_{i} L(y_i, f_{T-1} + h_m(x_i)) \qquad f_{T-1} = \hat{y}_{T-1}(x_i)$$

Expanding the Loss-function to the second order via Taylor:

$$L_T = \sum_{i} L(y_i, y_{T-1}) + \frac{dL}{d\hat{y}} h_m(x_i) + \frac{1}{2} \frac{d^2L}{d^2 \hat{y}} h_m^2(x_i)$$

Neglecting constant terms and using a more compat notation:

$$L_T = \sum_{i} g_T h_m(x_i) + \frac{1}{2} l_T h_m^2(x_i)$$

Extreme Boosting tree

$$L_T = \sum_{i} g_T h_m(x_i) + \frac{1}{2} l_T h_m^2(x_i)$$

Suppose first that we already have a partition into M regions R_1^* , ... R_M^* hence:

$$h_m(x) = \sum_{m=1}^{M} w_i I(x \in R_i) \qquad \frac{d}{dw_k} L_T = \sum_{i \in R_k^*} g_T + w_k l_T = 0 \rightarrow w_k^* = \frac{\sum_{i \in R_k^*} g_T(x_i)}{\sum_{i \in R_k^*} l_T(x_i)}$$

The coeff w_k compare only In the sum for the observation $i \in R_k^*$

Then use CART algorithm to build the tree based on loss L_T and optimal weight w_k^*

Extreme Boosting tree: alternative alg.

$$L_T = \sum_i g_t h_m(x_i) + \frac{1}{2} l_t h_m^2(x_i) \quad \text{Neglecting constant terms}$$

$$L_T = \sum_i \frac{1}{2} l_t(x_i) \left(-\frac{g_t}{l_t} - h_m(x_i) \right)^2$$

- f_o is a constant
- For each observation x_i compute $g_i = \frac{d}{df_0} L(y, f_0)|_{f_0}$ and $l_i = \frac{d^2}{d^2 f_0} L(y, f_0)|_{f_0}$
- Fit a regression tree $h_1(x; \gamma_1)$ on the dataset $\left\{x_i, -\frac{g_t}{l_t}\right\}$ minimizing $L_T = \sum_i \frac{1}{2} l_t(x_i) \left(-\frac{g_t}{l_t} h_m(x_i)\right)^2$
- Update $f_1(x) = f_0(x) + \epsilon f_1(x)$ where ϵ is an input parameter
- Iterate until $m \to M$

$$L = (y_i - f(x))^2$$

$$\frac{d}{df_0} L(y, f_0)|_{f_0} = (y_i - f_0)$$

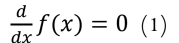
$$\frac{d^2}{d^2 f_0} L(y, f_0)|_{f_0} = -1$$

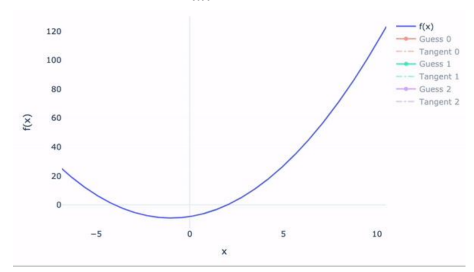
Extreme Boosting and Newton methods

$$L_T = \sum_{i} \frac{1}{2} l_t(x_i) \left(-\frac{g_t}{l_t} - h_m(x_i) \right)^2$$

...why the term $-\frac{g_t}{l_t}$ is not a surprise?

Suppose we want to solve a minimization problem (equivalent to root finding):





Newton Raphson Method:

$$x_0 = x_0$$

$$x_{t+1} = x_t - \frac{\frac{d}{dx}f(x)|_{x_t}}{\frac{d^2}{d^2x}f(x)|_{x_t}}$$

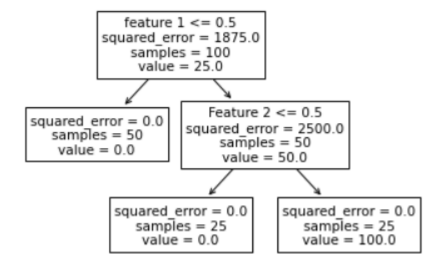
At convergence x_t is a solution of (1)

More on variable importance

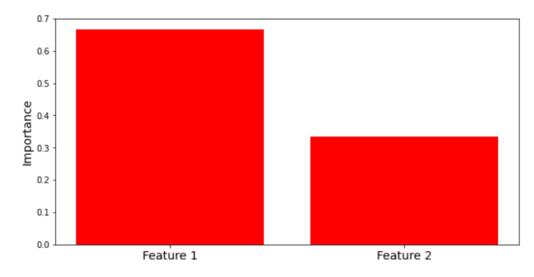
Inconsistency of variable importance

feature attribution methods introduce above are often inconsistent, meaning they can assign higher importance to features with a lower impact on the model's output.

 $y = (feature_1 \& feature_2) * 100$



We expect that feature 1 and feature 2 have same importance but..



Result of model.feature_importances_

Standard importance attribution methods gives an average picture of the contribution of the features, but we are more interested in understanding why a model made a prediction on a given record

SHAP method

IDEA: You do not understand the importance of the variables until you do not have it

Let us introduce a simpler explanation model g(z) used to explain the more complex model f(x)

$$g(z) = \phi_0 + \sum_P \phi_p z_p$$

 $z \in \{0,1\}^M$ where M is the total number of features. It represents a feature being observed $(z_i = 1)$ or unknown $(z_i = 0)$

 ϕ_i are the features importance values

We want that given an observation $\{x_i\} \to f(x_i) = g(z = [1, ... 1])$. In other word ϕ_i are a function of $\{x_i\}$

SHAP method

How to compute ϕ_i ? Using Shap-values from game theory!

All the possible model with the set S

Model considering the set S without the feature i

$$\phi_i = \sum_{S \le M - \{i\}} \frac{|S|! (|M| - |S| - 1)!}{|M|!} (f_{S + \{i\}}(x_i) - f_S(x_i))$$

S are the subsets of all M features

Model considering the set S and the feature i

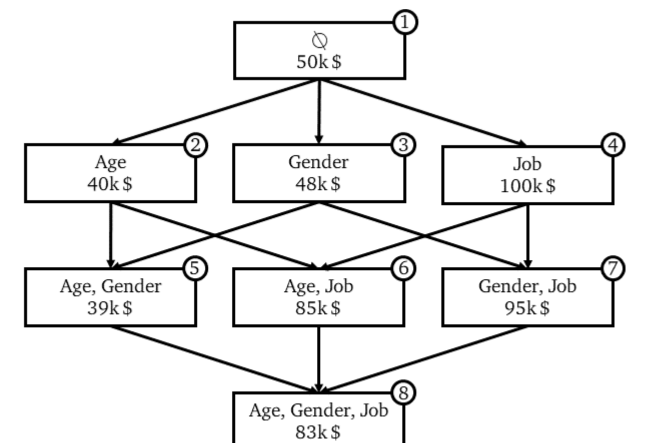
 ϕ_0 is the value predicted by the model without features, for regression is the average of y

Naive Implementation: Fit 2^M model with all possible subset from the set of features and then compute ϕ_i

Real Implementation: For tree model shap values can be computed exactly (why?), for other model there are some approximations!!

SHAP method: example

 $M = \{age, gender, job\}$



We want the importance of feature the age!

	without {age}	With {age}	difference	weight
Ø	50	40	-10	$\frac{2!}{3!} = \frac{1}{3}$
{Job}	100	85	-15	$\frac{1!}{3!} = \frac{1}{6}$
{Gender}	48	39	- 9	$\frac{1!}{3!} = \frac{1}{6}$
{Job, Gender}	95	83	-12	$\frac{2!}{3!} = \frac{1}{3}$

$$\phi_{age} = -11.33$$

f = 3

f = 0

f = 1

f = 2

SHAP method for classification

SHAP algorithm computes the SHAP values with respect to the margin not the transformed probability, the values you obtain are **log odds values!**

$$\log \frac{p}{1-p} = g(z) = \phi_0 + \sum_{p} \phi_p z_p$$