ML Methods
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(2) A Practical View
- Method or Models
- Interpretability
- Metric Choice
(3) A Better Point of View
- The Example of Univariate Linear Regression
- Supervised Learning

4) Risk Estimation and Method Choice

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Machine Learning



## The classical definition of Tom Mitchell

A computer program is said to learn from experience $\mathbf{E}$ with respect to some class of tasks $\mathbf{T}$ 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


## Article Clustering



An article clustering algorithm:

- Task: group articles corresponding to the same news
- Performance: quality of the clusters
- Experience: set of articles


## Smart Grid Controler



A controler in its sensors in a home smart grid:

- Task: control the devices in real-time
- Performance: energy costs
- Experience:
- previous days
- current environment and performed actions


Supervised and Unsupervised


## Supervised Learning (Imitation)

- Goal: Learn a function $f$ predicting a variable $Y$ from an individual $\underline{X}$.
- Data: Learning set with labeled examples $\left(\underline{X}_{i}, Y_{i}\right)$
- Assumption: Future data behaves as past data!
- Predicting is not explaining!

Supervised and Unsupervised


## Supervised Learning (Imitation)

- Goal: Learn a function $f$ predicting a variable $Y$ from an individual $X$.
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- Predicting is not explaining!

Unsupervised Learning (Structure Discovery)

- Goal: Discover a structure within a set of individuals $\left(X_{i}\right)$.
- Data: Learning set with unlabeled examples $\left(\underline{X}_{i}\right)$
- Unsupervised learning is not a well-posed setting. . .


## Machine Can and Cannot



## Machine Can

- Forecast (Prediction using the past)
- Detect expected changes
- Memorize/Reproduce/Imitate
- Take decisions very quickly
- Generate a lot of variations
- Learn from huge dataset
- Optimize a single task
- Help (or replace) some human beings


## Machine Cannot

- Predict something never seen before
- Detect any new behaviour
- Create something brand new
- Understand the world
- Plan by reasoning
- Get smart really fast
- Go beyond their task
- Replace (or kill) all human beings
- A lot of progresses but still very far from the singularity...


## Machine Learning



## 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.


## Machine Learning Pipeline



## Learning pipeline

- Test and compare models.
- Deployment pipeline is different!

Data Science $\neq$ 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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## Monthly KPI Dashboard

- Using financial data to display important KPI for top managers every month in a slide
- Automation to guaranty the quality of the results.



## Realtime Log Dashboard

- Use log data to show the state of a system to IT in real time using on-premise tools.
- Automation to handle the huge volumetry.


On-demand Legal Document Generation

- Use raw data to legal document template for a lawyer on-demand using a local database.
- First draft to be edited by the lawyer.

AB Testing


ER Waiting Time Prediction


## Real-Time ER Waiting Time Prediction

- Use patient data to provide in real time an estimate of the remaining waiting time to the ER patient.
- Tool helping to bear the wait.


Weekly Churn Prediction

- Using consumer characteristics and history to give a churn score to the marketing every week using the cloud.
- Automation to scale to the volumetry but no strategy recommendation.



## Realtime Automatic Fruit Sorting

- Using camera to sort fruits in a plant in realtime using local computers with GPU.
- Automation to reduce cost.


## Realtime Chatbot



## Realtime Chatbot

- Use previous interactions to predict answer to a consumer question in real time using the cloud.
- Reduce human interaction cost.



## Writing Assistant

- Enhance a text using AI in a communication system.
- Ease writing steps.



## Customer Segmentation

Data Science Project - Customer Segmentation

## Customer Segmentation

- Use customer data to suggest homogeneous groups to the marketing each year.
- Easier to think in term of groups than individuals


## Realtime Anomaly Detection



## Realtime Anomaly Detection

- Use production data to detect $2 n o m a l i e s ~ i n ~ a ~ p l a n t i n ~ r e a l ~ t i m e ~ o n ~ a ~ S c a d a ~ s y s t e m . ~$
- Reduce failure cost.


On-demand Fraud Detection

- Use claim and client data ty detect fraud for an insurer on-demand using on-premise resources
- First automated pass on the claims.


Prescriptive Maintenance (Not yet available. . .)

- Use data to devise and apply the best maintenance plan in a plant using IOT.
- Reduce maintenance cost.
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## What is a Method?



A Standard Machine Learning Pipeline

$B(x) \geq y$

## 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!

Simple Approach: Similarity


## 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


$$
\begin{aligned}
& Y_{i} a x_{+} b \\
& \sum\left|Y_{i}-\left(a x_{i}+4\right)\right|^{2} \\
& \sum\left|Y_{i}-\left(a x_{i}+3\right)\right|
\end{aligned}
$$

## Linear Method

- Simple formula: $a_{0}+a_{1} X^{(1)}+\cdots+a_{d} X^{(d)}$
- 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



## 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

Combing Simple Things: Ensemble


## 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

Chain Simple Things: Deep Learning


## 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

| Method | Performance | Training Speed | Inf. Speed | Stability | Interpretability |
| :--- | :---: | :---: | :---: | :---: | :---: |
| Similarity | - | $\emptyset$ | - | + | + |
| Linear | + | ++ | ++ | ++ | + |
| Tree | - | ++ | ++ | - | ++ |
| Ensemble | ++ | - | + | ++ | - |
| Deep | ++ | - | - | - | - |

## 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


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.



## ML Pipeline



## Learning pipeline

- Test and compare models.
- Deployment pipeline is different!


## Cross Validation Principle



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

- Check the quality of a method by repeating the previous approach.
- Beware: a different predictor is learnt for each split.

The Full Cross Validation Scheme
$\square$



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



## Competition results

- The true model is not the winner!


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Interpretation?


## Is this that easy?

- Simple formula setting:

$$
Y \simeq f(X)=a_{0}+a_{1} X^{(1)}+a_{2} X^{(2)}+\cdots+a_{d} X^{(d)}
$$

- Beware of the interpretation!
- Everything being equal. . . Correlation is not causality. . .



## Intepretability 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.


## eXplainable AI (XAI)



## A few directions

- Data Explaination.
- Use of explainable methods (linear?).
- Use of black box methods:
- Global explanation (variable importance)
- Local explanation (linear approximationn, alternative scenario...)
- Causality very hard to access without a real experimental plan with interventions!


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## Metric and Solution

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.



## 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, idealy encoding domain expertise (asymmetry...)
- The criterion can be different between optimization and evaluation because of computation requirements.
- Very important factor (too) often neglicted.



## 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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## Eucalyptus



- Simple (and classical) dataset.
- Goal: predict the height from circumference
- $X=$ circ $=$ circumference.
- $Y=h t=$ height.


## Eucalyptus



## Linear Model

- Parametric model:

$$
f_{\beta}(\operatorname{circ})=\beta^{(1)}+\beta^{(2)} \operatorname{circ}
$$

- How to choose $\beta=\left(\beta^{(1)}, \beta^{(2)}\right)$ ?


## Methodology

- Natural goodness criterion:

$$
\begin{aligned}
\sum_{i=1}^{n}\left|Y_{i}-f_{\beta}\left(\underline{X}_{i}\right)\right|^{2} & =\sum_{i=1}^{n}\left|h t_{i}-f_{\beta}\left(\operatorname{circ}_{i}\right)\right|^{2} \\
& =\sum_{i=1}^{n}\left|h t_{i}-\left(\beta^{(1)}+\beta^{(2)} \operatorname{circ}_{i}\right)\right|^{2}
\end{aligned}
$$

- Choice of $\beta$ that minimizes this criterion!

$$
\widehat{\beta}=\underset{\beta \in \mathbb{R}^{2}}{\operatorname{argmin}} \sum_{i=1}^{n}\left|h_{i}-\left(\beta^{(1)}+\beta^{(2)} \operatorname{circ}_{i}\right)\right|^{2}
$$

- Easy minimization with an explicit solution!

$$
y=\frac{(1 x)}{x} \beta+\varepsilon \quad \hat{\beta}=\left[(1 x)^{\top}(1 x)\right]^{-L}(1 x)^{\top} y
$$



## Prediction

- Linear prediction for the height:

$$
\widehat{\mathrm{ht}}=f_{\widehat{\beta}}(\operatorname{circ})=\widehat{\beta}^{(1)}+\widehat{\beta}^{(2)} \operatorname{circ}
$$

## Linear Regression

- Statistical model: $\left(\operatorname{circ}_{i}, \mathrm{ht}_{i}\right)$ i.i.d. with the same law as a generic (circ, ht).
- Performance criterion: Look for $f$ with a small average error

$$
\mathbb{E}\left[|\mathrm{ht}-f(\operatorname{circ})|^{2}\right]
$$

- Empirical criterion: Replace the unknown law by its empirical counterpart

$$
\frac{1}{n} \sum_{i=1}^{n}\left|\mathrm{ht}_{i}-f\left(\operatorname{circ}_{i}\right)\right|^{2}
$$

- Predictor model: As the minimum over all function is 0 (if all the $\mathrm{circ}_{i}$ are different), restrict to the linear functions $f(\operatorname{circ})=\beta^{(1)}+\beta^{(2)}$ circ to avoid over-fitting.
- Model fitting: Explicit formula here.
- This model can be too simple!



## Polynomial Model

- Polynomial model: $f_{\beta}(\operatorname{circ})=\sum_{l=1}^{p} \beta^{(I)} \operatorname{circ}^{I-1}$
- 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

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## Which Degree?



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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?


Model complexity

## 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 fit a model to the data?
- How to assess its quality?
- How to select a model among a collection? Volidalen
Estinulen de lo qualité
- 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 $Y \in \mathcal{Y}$.
- $(\underline{X}, Y) \sim \mathbb{P}$ with $\mathbb{P}$ unknown.
- Training data: $\mathcal{D}_{n}=\left\{\left(\underline{X}_{1}, Y_{1}\right), \ldots,\left(\underline{X}_{n}, Y_{n}\right)\right\} \quad$ (i.i.d. $\left.\sim \mathbb{P}\right)$
- Often
- $\underline{X} \in \mathbb{R}^{d}$ and $Y \in\{-1,1\}$ (classification) $<Y \in\{1,2,3, \ldots \in\}$
- or $\underline{X} \in \mathbb{R}^{d}$ and $Y \in \mathbb{R}$ (regression).
- A predictor is a function in $\mathcal{F}=\{f: \mathcal{X} \rightarrow \mathcal{Y}$ meas. $\}$


## Goal

- Construct a good predictor $\widehat{f}$ from the training data.
- Need to specify the meaning of good.
- Classification and regression are almost the same problem!


## Loss function for a generic predictor

- Loss function: $\ell(Y, f(\underline{X}))$ measures the goodness of the prediction of $Y$ by $f(\underline{X})$
- Examples:
- $0 / 1$ loss: $\ell(Y, f(\underline{X}))=\mathbf{1}_{Y \neq f(X)}$
- Quadratic loss: $\ell(Y, f(\underline{X}))=|Y-f(\underline{X})|^{2}$


## Risk function

- Risk measured as the average loss for a new couple:

$$
\mathcal{R}(f)=\mathbb{E}_{(X, Y) \sim \mathbb{P}}[\ell(Y, f(\underline{X}))]
$$

- Examples:
- $0 / 1$ loss: $\mathbb{E}[\ell(Y, f(\underline{X}))]=\mathbb{P}(Y \neq f(\underline{X}))$
- Quadratic loss: $\mathbb{E}[\ell(Y, f(\underline{X}))]=\mathbb{E}\left[|Y-f(\underline{X})|^{2}\right]$
- Beware: As $\widehat{f}$ depends on $\mathcal{D}_{n}, \mathcal{R}(\widehat{f})$ is a random variable!
- The best solution $f^{\star}$ (which is independent of $\mathcal{D}_{n}$ ) is

$$
f^{\star}=\arg \min _{f \in \mathcal{F}} \mathcal{R}(f)=\arg \min _{f \in \mathcal{F}} \mathbb{E}[\ell(Y, f(\underline{X}))]=\arg \min _{f \in \mathcal{F}} \mathbb{E}_{\underline{X}}\left[\mathbb{E}_{Y \mid \underline{X}}[\ell(Y, f(\underline{X}))]\right]
$$

## Bayes Predictor (explicit solution)

- In binary classification with $0-1$ loss:

$$
f^{\star}(\underline{X})= \begin{cases}+1 & \text { if } \mathbb{P}(Y=+1 \mid \underline{X}) \geq \mathbb{P}(Y=-1 \mid \underline{X}) \\ & \Leftrightarrow \mathbb{P}(Y=+1 \mid \underline{X}) \geq 1 / 2 \\ -1 & \text { otherwise }\end{cases}
$$

- In regression with the quadratic loss

$$
f^{\star}(\underline{X})=\mathbb{E}[Y \mid \underline{X}]
$$

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

## Machine Learning

- Learn a rule to construct a predictor $\widehat{f} \in \mathcal{F}$ from the training data $\mathcal{D}_{n}$ s.t. the risk $\mathcal{R}(\widehat{f})$ 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 $\mathcal{S}=\left\{f_{\theta}, \theta \in \Theta\right\}$
- One replaces the minimization of the average loss by the minimization of the empirical loss

$$
\widehat{f}=f_{\widehat{\theta}}=\underset{f_{\theta}, \theta \in \Theta}{\operatorname{argmin}} \frac{1}{n} \sum_{i=1}^{n} \ell\left(Y_{i}, f_{\theta}\left(\underline{X}_{i}\right)\right)
$$

- Examples:
- Linear regression
- Linear classification with

$$
\mathcal{S}=\left\{\underline{x} \mapsto \operatorname{sign}\left\{\underline{x}^{\top} \beta+\beta^{(0)}\right\} / \beta \in \mathbb{R}^{d}, \beta^{(0)} \in \mathbb{R}\right\}
$$

## 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.


Logistic


Naive Bayes with kernel density estimates


Eucalyptus


## Dataset - P.A. Cornillon

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

Eucalyptus



Eucalyptus


Eucalyptus


## Under-fitting / Over-fitting Issue



## Model Complexity Dilemna

- What is best a simple or a complex model?
- Too simple to be good? Too complex to be learned?



## Under-fitting / Over-fitting

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


## Bias-Variance Dilemma

- General setting:
- $\mathcal{F}=\{$ measurable functions $\mathcal{X} \rightarrow \mathcal{Y}\}$
- Best solution: $f^{\star}=\operatorname{argmin}_{f \in \mathcal{F}} \mathcal{R}(f)$
- Class $\mathcal{S} \subset \mathcal{F}$ of functions
- Ideal target in $\mathcal{S}: f_{\mathcal{S}}^{\star}=\operatorname{argmin}_{f \in \mathcal{S}} \mathcal{R}(f)$
- Estimate in $\mathcal{S}: \widehat{f}_{\mathcal{S}}$ obtained with some procedure


Approximation error and estimation error (Bias/Variance)

$$
\mathcal{R}\left(\widehat{f}_{\mathcal{S}}\right)-\mathcal{R}\left(f^{\star}\right)=\underbrace{\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right)-\mathcal{R}\left(f^{\star}\right)}_{\text {Approximation error }}+\underbrace{\mathcal{R}\left(\widehat{f}_{\mathcal{S}}\right)-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right)}_{\text {Estimation error }}
$$

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


## Agnostic approach

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

Under-fitting / Over-fitting Issue


- 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 $\Longleftrightarrow$ avoid overfitting and underfitting

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


## Statistical Learning Analysis

- Error decomposition:

$$
\mathcal{R}\left(\widehat{f}_{\mathcal{S}}\right)-\mathcal{R}\left(f^{\star}\right)=\underbrace{\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right)-\mathcal{R}\left(f^{\star}\right)}_{\text {Approximation error }}+\underbrace{\mathcal{R}\left(\widehat{f}_{\mathcal{S}}\right)-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right)}_{\text {Estimation error }}
$$

- 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 $\mathbb{P} \ldots$ (Nonparametric Statistics?)


$$
\begin{aligned}
& f: x \rightarrow\{-i x \\
& b: x \rightarrow \mathbb{R} \\
& \downarrow x y m \\
&\{-1,1\}
\end{aligned}
$$

## Empirical Risk Minimizer

$$
\widehat{f}=\underset{f \in \mathcal{S}}{\operatorname{argmin}} \frac{1}{n} \sum_{i=1}^{n} \ell^{0 / 1}\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)
$$

- Classification loss: $\ell^{0 / 1}(y, f(\underline{x}))=\mathbf{1}_{y \neq f(\underline{x})}$
- Not convex and not smooth!


## Probabilistic Point of View

## Ideal Solution and Estimation



- The best solution $f^{\star}$ (which is independent of $\mathcal{D}_{n}$ ) is

$$
f^{\star}=\arg \min _{f \in \mathcal{F}} \mathcal{R}(f)=\arg \min _{f \in \mathcal{F}} \mathbb{E}[\ell(Y, f(\underline{X}))]=\arg \min _{f \in \mathcal{F}} \mathbb{E}_{\underline{X}}\left[\mathbb{E}_{Y \mid \underline{X}}[\ell(Y, f(\underline{x}))]\right]
$$

## Bayes Predictor (explicit solution)

In binary classification with $0-1$ loss:

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

- Issue: Solution requires to know $\mathbb{E}[Y \mid \underline{X}]$ for all values of $\underline{X}$ !
- Solution: Replace it by an estimate.

Optimization Point of View
Loss Convexification


Minimizer of the risk

$$
\widehat{f}=\underset{f \in \mathcal{S}}{\operatorname{argmin}} \frac{1}{n} \sum_{i=1}^{n} \ell^{0 / 1}\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)
$$

- 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

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

Canonical approach: $\widehat{f}_{\mathcal{S}}=\operatorname{argmin}_{f \in \mathcal{S}} \frac{1}{n} \sum_{i=1}^{n} \ell\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)$

## Problems

- How to choose $\mathcal{S}$ ?
- How to compute the minimization?


## A Probabilistic Point of View

Solution: For $\underline{X}$, estimate $Y \mid \underline{X}$ 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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- Metric Choice


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- Two classes.
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# Example: Linear Discrimination 

Logistic


Naive Bayes with kernel density estimates

k-NN with $\mathrm{k}=1$

k-NN with $\mathrm{k}=5$


k-NN with k=9
Decision region


Decision boundary

$\mathrm{k}-\mathrm{NN}$ with $\mathrm{k}=13$
Decision region


Decision boundary

$\mathrm{k}-\mathrm{NN}$ with $\mathrm{k}=17$
Decision region


Decision boundary

k-NN with $\mathrm{k}=21$

k-NN with $\mathrm{k}=25$
Decision region


Decision boundary

k-NN with $\mathrm{k}=29$
Decision region


Decision boundary

k-NN with $\mathrm{k}=33$
Decision region


Decision boundary

k-NN with $\mathrm{k}=37$
Decision region


Decision boundary

$\mathrm{k}-\mathrm{NN}$ with $\mathrm{k}=45$
Decision region


Decision boundary

k-NN with $\mathrm{k}=53$
Decision region


Decision boundary

k-NN with $\mathrm{k}=61$
Decision region


Decision boundary

k-NN with $\mathrm{k}=69$
Decision region


Decision boundary

k-NN with $\mathrm{k}=77$
Decision region


Decision boundary

k-NN with $\mathrm{k}=85$
Decision region


Decision boundary

$\mathrm{k}-\mathrm{NN}$ with $\mathrm{k}=101$
Decision region


Decision boundary

k-NN with $\mathrm{k}=109$
Decision region


Decision boundary

$k-N N$ with $\mathrm{k}=117$
Decision region


Decision boundary

k -NN with $\mathrm{k}=125$

k-NN with $\mathrm{k}=133$
Decision region


Decision boundary

$\mathrm{k}-\mathrm{NN}$ with $\mathrm{k}=141$

$\mathrm{k}-\mathrm{NN}$ with $\mathrm{k}=149$

k-NN with $\mathrm{k}=157$
Decision region


Decision boundary

k-NN with $\mathrm{k}=165$
Decision region


Decision boundary

k-NN with $\mathrm{k}=173$
Decision region


Decision boundary

k-NN with $\mathrm{k}=181$
Decision region


Decision boundary

k-NN with $\mathrm{k}=189$
Decision region


Decision boundary

k-NN with $\mathrm{k}=197$
Decision region


Decision boundary


## Training Risk Issue

Model complexity

## Risk behaviour

- 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 $\mathcal{S}$

$$
R_{n}\left(\widehat{f_{\mathcal{S}}}\right) \rightarrow R_{n}\left(\widehat{f_{\mathcal{S}}}\right)+\operatorname{cor}(\mathcal{S})
$$

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.

Cross Validation


- 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,

$$
\begin{aligned}
& \frac{1}{n} \sum l\left(x_{i}, f(x)\right) \simeq \mathbb{E}[l(1, l(x))] \\
& \frac{1}{i} \sum l\left(y_{i}, \hat{l}(x)\right)=\mathbb{E}\left[l\left(y_{1} \hat{l}(x)\right]\right.
\end{aligned}
$$

- $V$-fold cross validation.

$$
\begin{aligned}
& {\underset{\alpha}{n}}_{\mathbb{E}}^{[ } \frac{1}{n}\left[l\left(y_{1}, \hat{l}(x)\right)\right] \\
& \leq \mathbb{E}_{c_{0}}\left[\hat{n}_{n}^{n} l\left(\frac{1}{1}, l^{x}\left(x_{1}\right)\right]\right. \\
& =\mathbb{E}\left[l\left(1 /, l^{*}(x)\right) 100\right.
\end{aligned}
$$

## 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 $\widehat{f}^{H O}$ from the subset $\mathcal{D}_{\text {train }}$.
- Compute the empirical risk on the subset $\mathcal{D}_{\text {test }}$ :

$$
\mathcal{R}_{n}^{H O}\left(\widehat{f}^{H O}\right)=\frac{1}{n \epsilon} \sum_{\left(\underline{X}_{i}, Y_{i}\right) \in \mathcal{D}_{\text {test }}} \ell\left(Y_{i}, \widehat{f}^{H O}\left(\underline{X}_{i}\right)\right)
$$

## Predictor Risk Estimation

- Use $\hat{f}^{H O}$ as predictor.
- Use $\mathcal{R}_{n}^{H O}\left(\hat{f}^{H O}\right)$ as an estimate of the risk of this estimator.


## Method Selection by Cross Validation

- Compute $\mathcal{R}_{n}^{H O}\left(\widehat{f}_{\mathcal{S}}^{H O}\right)$ for all the considered methods,
- Select the method with the smallest CV risk,
- Reestimate the $\widehat{f}_{\mathcal{S}}$ with all the data.

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 $\widehat{f}^{H O}$ from the subset $\mathcal{D}_{\text {train }}$.
- Compute the empirical risk on the subset $\mathcal{D}_{\text {test }}$ :

$$
\mathcal{R}_{n}^{H O}\left(\widehat{f}^{H O}\right)=\frac{1}{n \epsilon} \sum_{\left(\underline{X}_{i}, Y_{i}\right) \in \mathcal{D}_{\text {test }}} \ell\left(Y_{i}, \widehat{f}^{H O}\left(\underline{X}_{i}\right)\right)
$$

- 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 $\mathcal{R}_{n}^{\mathrm{HO}}\left(\hat{f}^{\mathrm{HO}}\right)$ not taken into account.
$V$-fold Cross Validation


Principle

- Split the dataset $\mathcal{D}$ in $V$ sets $\mathcal{D}_{\vee}$ of almost equals size.
- For $v \in\{1, . ., V\}$ :
- Learn $\widehat{f}^{-v}$ from the dataset $\mathcal{D}$ minus the set $\mathcal{D}_{v}$.
- Compute the empirical risk:

$$
\mathcal{R}_{n}^{-v}\left(\widehat{f}^{-v}\right)=\frac{1}{n_{v}} \sum_{\left(X_{i}, Y_{i}\right) \in \mathcal{D}_{v}} \ell\left(Y_{i}, \widehat{f}^{-v}\left(\underline{X}_{i}\right)\right)
$$

- Compute the average empirical risk:

$$
\mathcal{R}_{n}^{C V}(\widehat{f})=\frac{1}{V} \sum_{v=1}^{V} \mathcal{R}_{n}^{-v}\left(\widehat{f}^{-v}\right)
$$

- Estimation of the quality of a method not of a given predictor.
- Leave One Out: V = n.


## $V$-fold Cross Validation

## Analysis (when $n$ is a multiple of $V$ )

- The $\mathcal{R}_{n}^{-v}\left(\widehat{f}^{-v}\right)$ are identically distributed variable but are not independent!
- Consequence:

$$
\begin{aligned}
\mathbb{E}\left[\mathcal{R}_{n}^{C V}(\widehat{f})\right]= & \mathbb{E}\left[\mathcal{R}_{n}^{-v}\left(\widehat{f}^{-v}\right)\right] \\
\operatorname{Var}\left[\mathcal{R}_{n}^{C V}(\widehat{f})\right]= & \frac{1}{V} \operatorname{Var}\left[\mathcal{R}_{n}^{-v}\left(\widehat{f}^{-v}\right)\right] \\
& +\left(1-\frac{1}{V}\right) \operatorname{Cov}\left[\mathcal{R}_{n}^{-v}\left(\widehat{f}^{-v}\right), \mathcal{R}_{n}^{-v^{\prime}}\left(\widehat{f}^{-v^{\prime}}\right)\right]
\end{aligned}
$$

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

$$
\frac{1}{\Delta+\frac{1}{v}}
$$

- Accuracy/Speed tradeoff: $V=5$ or $V=10 \ldots$
- 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,

$$
\widehat{f}^{-i}\left(\underline{X}_{i}\right)=\frac{\widehat{f}\left(X_{i}\right)-h_{i i} Y_{i}}{1-h_{i i}}
$$

with $h_{i i}$ the ith diagonal coefficient of the hat (projection) matrix.

- Proof based on linear algebra!
- Leads to a fast formula for LOO:

$$
\mathcal{R}_{n}^{L O O}(\widehat{f})=\frac{1}{n} \sum_{i=1}^{n} \frac{\left|Y_{i}-\widehat{f}\left(\underline{X}_{i}\right)\right|^{2}}{\left(1-h_{i i}\right)^{2}}
$$

# Cross Validation 

variable

- Accuracy
- AccuracyCV


# Example: $\operatorname{KNN}(\hat{k}=61$ using cross-validation) 

$\mathrm{k}-\mathrm{NN}$ with $\mathrm{k}=61$


## Train/Validation/Test



- 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!
- 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

$$
\underset{\theta \in \Theta}{\operatorname{argmin}} \frac{1}{n} \sum_{i=1}^{n} \ell\left(Y_{i}, f_{\theta}\left(\underline{X}_{i}\right)\right)+\operatorname{pen}(\theta)
$$

where $\operatorname{pen}(\theta)$ 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 pen $(\theta)=2 \frac{d}{n} \sigma^{2}$.
- AIC Heuristics: Maximum Likelihood with pen $(\theta)=\frac{d}{n}$.
- BIC Heuristics: Maximum Likelohood with $\operatorname{pen}(\theta)=\log (n) \frac{d}{n}$.
- Structural Risk Minimization: Pred. loss and clever penalty.


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## Comparison of Two Means

## Means

- Setting: r.v. $e_{i}^{(I)}$ with $1 \leq i \leq n_{I}$ and $I \in\{1,2\}$ and their means

$$
\overline{e^{(I)}}=\frac{1}{n_{l}} \sum_{i=1}^{n_{l}} e_{i}^{(l)}
$$

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


## Classical i.i.d setting

- Assumption: $e_{i}^{(I)}$ are i.i.d. for each $/$.
- Test formulation: Can we reject the null hypothesis that $\mathbb{E}\left[e^{(1)}\right]=\mathbb{E}\left[e^{(2)}\right]$ ?
- 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 / but not necessarily independent.
- Test formulation: Can we reject the null hypothesis that $\mathbb{E}\left[e^{(1)}\right]=\mathbb{E}\left[e^{(2)}\right]$ ?
- 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: $e_{i}^{(I)}$ are i.d. for each / but not necessarily independent.
- Tests formulation:
- Can we reject the null hypothesis that the $\mathbb{E}\left[e^{(I)}\right]$ 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.



## 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.


## Probabibly-Approximately-Correct (PAC) Approach

- Replace the control on the average risk by a probabilistic bound

$$
\mathbb{P}(\mathbb{E}[\ell(Y, \hat{f}(\underline{X}))]>R) \leq \epsilon
$$

- 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 ~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, ~ 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 



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# Unbalanced and Rebalanced Dataset 



## Unbalanced Class

- Setting: One of the classes 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

- 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!


## Testing

- Testing class prob.: $\pi_{t}(k)$
- Testing risk target:

$$
\begin{aligned}
& \mathbb{E}_{\pi_{t}}[\ell(Y, f(\underline{X}))]= \\
& \sum_{k} \pi_{t}(k) \mathbb{E}[\ell(Y, f(\underline{X})) \mid Y=k]
\end{aligned}
$$

## Training

- Training class prob.: $\pi_{t r}(k)$
- Training risk target:

$$
\begin{aligned}
& \mathbb{E}_{\pi_{t r}}[\ell(Y, f(\underline{X}))]= \\
& \sum_{k} \pi_{t r}(k) \mathbb{E}[\ell(Y, f(\underline{X})) \mid Y=k]
\end{aligned}
$$

## Implicit Testing Risk Using the Training One

- Amounts to use a weighted loss:

$$
\begin{aligned}
\mathbb{E}_{\pi_{t r}}[\ell(Y, f(\underline{X}))]= & \sum_{k} \pi_{t r}(k) \mathbb{E}[\ell(Y, f(\underline{X})) \mid Y=k] \\
& =\sum_{k} \pi_{t}(k) \mathbb{E}\left[\left.\frac{\pi_{t r}(k)}{\pi_{t}(k)} \ell(Y, f(\underline{X})) \right\rvert\, Y=k\right] \\
& =\mathbb{E}_{\pi_{t}}\left[\frac{\pi_{t r}(Y)}{\pi_{t}(Y)} \ell(Y, f(\underline{X}))\right]
\end{aligned}
$$

- 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:

$$
\ell(Y, f(\underline{X})) \rightarrow C(Y) \ell(Y, f(\underline{X}))
$$

- Weighted risk target:

$$
\mathbb{E}[C(Y) \ell(Y, f(\underline{X}))]
$$

- Rk: Strong link with $\ell$ as $C$ is independent of $f$.
- Often allow reusing algorithm constructed for $\ell$.
- C may also depend on $X \ldots$
- The Bayes classifier is now:

$$
f^{\star}=\operatorname{argmin} \mathbb{E}[C(Y) \ell(Y, f(\underline{X}))]=\operatorname{argmin} \mathbb{E}_{\underline{X}}\left[\mathbb{E}_{Y \mid \underline{X}}[C(Y) \ell(Y, f(\underline{X}))]\right]
$$

## Bayes Predictor

- For $\ell^{0 / 1}$ loss,

$$
f^{\star}(\underline{X})=\underset{k}{\operatorname{argmax}} C(k) \mathbb{P}(Y=k \mid \underline{X})
$$

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


## Cost and Proportions

- Testing risk target:

$$
\mathbb{E}_{\pi_{t}}\left[C_{t}(Y) \ell(Y, f(\underline{X}))\right]=\sum_{k} \pi_{t}(k) C_{t}(k) \mathbb{E}[\ell(Y, f(\underline{X})) \mid Y=k]
$$

- Training risk target

$$
\mathbb{E}_{\pi_{t r}}\left[C_{t r}(Y) \ell(Y, f(\underline{X}))\right]=\sum_{k} \pi_{t r}(k) C_{t r}(k) \mathbb{E}[\ell(Y, f(\underline{X})) \mid Y=k]
$$

- Coincide if

$$
\pi_{t}(k) C_{t}(k)=\pi_{t r}(k) C_{t r}(k)
$$

- Lots of flexibility in the choice of $C_{t}, C_{t r}$ or $\pi_{t r}$.

Weighted Loss and Resampling

- Weighted loss: choice of a weight $C_{t} \neq 1$.
- Resampling: use a $\pi_{t r} \neq \pi_{t}$.
- Stratified sampling may be used to reduce the size of a dataset without loosing a low probability class!


## Combining Weights and Resampling

- Weighted loss: use $C_{t r}=C_{t}$ as $\pi_{t r}=\pi_{t}$.
- Resampling: use an implicit $C_{t}(k)=\pi_{t r}(k) / \pi_{t}(k)$.
- Combined: use $C_{t r}(k)=C_{t}(k) \pi_{t}(k) / \pi_{t r}(k)$
- Most ML methods allow such weights!


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A Probabilistic Point of View- Machine Learning- Motivation
(2)

A Practical View

- Method or Models
- Interpretability
- Metric Choice
(3)

A Better Point of View

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## Auto ML



## 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



## Auto ML Task

- Input:
- a dataset $\mathcal{D}=\left(\underline{X}_{i}, Y_{i}\right)$
- a loss function $\ell(Y, f(\underline{X}))$
- a set of possible predictors $f_{l, h, \theta}$ corresponding to a method / in a list, with hyperparameters $h$ and parameters $\theta$
- Output:
- a predictor $f$ equal to $f_{\hat{i}, \hat{h}, \hat{\theta}}$ 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



## 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

- Almost always (slightly) 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 $f_{\theta}(\underline{X})=\underline{X}^{\top} \beta+\beta^{(0)}$ with $\theta=\left(\beta, \beta^{(0)}\right)$.
- Let $\mathbb{P}_{\theta}(Y=1 \mid \underline{X})=e^{+f_{\theta}(\underline{X})} /\left(1+e^{f_{\theta}(\underline{X})}\right)$
- Estimate $\theta$ by $\hat{\theta}$ using a Maximum Likelihood.
- Classify using $\mathbb{P}_{\hat{\theta}}(Y=1 \mid \underline{X})>1 / 2$


## k Nearest Neighbors

- For any $\underline{X}^{\prime}$, define $\mathcal{V}_{\underline{X}^{\prime}}$ as the $k$ closest samples $X_{i}$ from the dataset.
- Compute a score $g_{k}=\sum_{X_{i} \in \mathcal{V}_{\underline{x}^{\prime}}} \mathbf{1}_{Y_{i}=k}$
- Classify using arg $\max g_{k}$ (majority vote).


## 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 $\ln (\mathbb{P}(\underline{X} \mid Y=k))+\ln (\mathbb{P}(Y=k))$

$$
\begin{aligned}
g_{k}(\underline{X})=- & \frac{1}{2}\left(\underline{X}-\mu_{k}\right)^{\top} \Sigma_{k}^{-1}\left(\underline{X}-\mu_{k}\right) \\
& -\frac{d}{2} \ln (2 \pi)-\frac{1}{2} \ln \left(\left|\Sigma_{k}\right|\right)+\ln (\mathbb{P}(Y=k))
\end{aligned}
$$

- Classify using arg max $g_{k}$
- Those three methods rely on a similar heuristic: the probabilistic point of view!
- The best solution $f^{\star}$ (which is independent of $\mathcal{D}_{n}$ ) is

$$
f^{\star}=\arg \min _{f \in \mathcal{F}} R(f)=\arg \min _{f \in \mathcal{F}} \mathbb{E}[\ell(Y, f(\underline{X}))]=\arg \min _{f \in \mathcal{F}} \mathbb{E}_{\underline{X}}\left[\mathbb{E}_{Y \mid \underline{X}}[\ell(Y, f(\underline{X})]]\right.
$$

## Bayes Predictor (explicit solution)

- In binary classification with $0-1$ loss:

$$
f^{\star}(\underline{X})= \begin{cases}+1 & \text { if } \mathbb{P}(Y=+1 \mid \underline{X}) \geq \mathbb{P}(Y=-1 \mid \underline{X}) \\ & \Leftrightarrow \mathbb{P}(Y=+1 \mid \underline{X}) \geq 1 / 2 \\ -1 & \text { otherwise }\end{cases}
$$

- In regression with the quadratic loss

$$
f^{\star}(\underline{X})=\mathbb{E}[Y \mid \underline{X}]
$$

Issue: Explicit solution requires to know $Y \mid \underline{X}($ or $\mathbb{E}[Y \mid \underline{X}])$ for all values of $X$ !

- Idea: Estimate $Y \mid \underline{X}$ by $\widehat{Y \mid \underline{X}}$ and plug it the Bayes classifier.


## Plugin Bayes Predictor

- In binary classification with $0-1$ loss:

$$
\widehat{f}(\underline{X})= \begin{cases}+1 & \text { if } \overline{\mathbb{P}(Y=+1 \mid \underline{X})} \geq \overline{\mathbb{P}(Y=-1 \mid \underline{X})} \\ & \Leftrightarrow \overline{\mathbb{P}(Y=+1 \mid \underline{X})} \geq 1 / 2 \\ -1 & \text { otherwise }\end{cases}
$$

- In regression with the quadratic loss

$$
\widehat{f}(\underline{X})=\mathbb{E}[\widehat{Y \mid \underline{X}}]
$$

- Rk: Direct estimation of $\mathbb{E}[Y \mid \underline{X}]$ by $\widehat{\mathbb{E}[Y \mid \underline{X}]}$ also possible...

Plugin Predictor

- How to estimate $Y \mid \underline{X}$ ?


## Three main heuristics

- Parametric Conditional modeling: Estimate the law of $Y \mid \underline{X}$ by a parametric law $\mathcal{L}_{\theta}(\underline{X})$ : (generalized) linear regression...
- Non Parametric Conditional modeling: Estimate the law of $Y \mid \underline{X}$ 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 $Y \mid \underline{X}: L D A / Q D A$, Naive Bayes...
- Rk: Direct estimation of $\mathbb{E}[Y \mid \underline{X}]$ by $\widehat{\mathbb{E}[Y \mid \underline{X}]}$ also possible...
- Input: a data set $\mathcal{D}_{n}$

Learn $Y \mid \underline{X}$ or equivalently $\mathbb{P}(Y=k \mid \underline{X})$ (using the data set) and plug this estimate in the Bayes classifier

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

$$
\widehat{f}(\underline{X})= \begin{cases}+1 & \text { if } \mathbb{P}(\widehat{Y=1} \mid \underline{X}) \geq \mathbb{P}(\widehat{Y=-1} \mid \underline{X}) \\ -1 & \text { otherwise }\end{cases}
$$

- Can we guaranty that the classifier is good if $Y \mid \underline{X}$ is well estimated?


## Classification Risk Analysis

## Theorem

- If $\widehat{f}=\operatorname{sign}\left(2 \widehat{p}_{+1}-1\right)$ then

$$
\begin{aligned}
\mathbb{E}\left[\ell^{0,1}(Y, \widehat{f}(\underline{X}))\right] & -\mathbb{E}\left[\ell^{0,1}\left(Y, f^{\star}(\underline{X})\right)\right] \\
& \leq \mathbb{E}\left[\|\widehat{Y \mid \underline{X}}-Y \mid \underline{X}\|_{1}\right] \\
& \leq\left(\mathbb{E}[2 \mathrm{KL}(Y \mid \underline{X}, \widehat{Y \mid \underline{X}}])^{1 / 2}\right.
\end{aligned}
$$

- If one estimates $\mathbb{P}(Y=1 \mid \underline{X})$ well then one estimates $f^{\star}$ 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 $\mathbb{P}(Y=1 \mid \underline{X})$ 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 $Y \mid \underline{X}$ by a parametric conditional density $\mathbb{P}_{\theta}(Y \mid \underline{X})$.


## Maximum Likelihood Approach

- Classical choice for $\theta$ :

$$
\widehat{\theta}=\underset{\theta}{\operatorname{argmin}}-\sum_{i=1}^{n} \log \mathbb{P}_{\theta}\left(Y_{i} \mid \underline{X}_{i}\right)
$$

- Goal: Minimize the Kullback-Leibler divergence between the conditional law of $Y \mid \underline{X}$ and $\mathbb{P}_{\theta}(Y \mid \underline{X})$

$$
\mathbb{E}\left[K L\left(Y \mid \underline{X}, \mathbb{P}_{\theta}(Y \mid \underline{X})\right)\right]
$$

- Rk: This is often not (exactly) the learning task!
- Large choice for the family $\left\{\mathbb{P}_{\theta}(Y \mid \underline{X})\right\}$ but depends on $\mathcal{Y}$ (and $\mathcal{X}$ ).
- Regression: One can also model directly $\mathbb{E}[Y \mid \underline{X}]$ by $f_{\theta}(\underline{X})$ and estimate it with a least-squares criterion. . .


## Linear Models

- Classical choice: $\theta=\left(\theta^{\prime}, \varphi\right)$

$$
\mathbb{P}_{\theta}(Y \mid \underline{X})=\mathbb{P}_{\underline{X}^{\top}{ }_{\beta, \varphi}}(Y)
$$

- Very strong modeling assumption!
- Classical examples:
- Binary variable: logistic, probit...
- Discrete variable: multinomial logistic regression...
- Integer variable: Poisson regression...
- Continuous variable: Gaussian regression...


## Binary Classifier <br> $$
R(y=1 \mid x)=h\left(\left\langle x_{1} \beta\right\rangle+\beta^{(2)}\right)
$$

## Plugin Linear Classification

- Model $\mathbb{P}(Y=+1 \mid \underline{X})$ by $h\left(\underline{X}^{\top} \beta+\beta^{(0)}\right)$ with $h$ non decreasing.
- $h\left(\underline{X}^{\top} \beta+\beta^{(0)}\right)>1 / 2 \Leftrightarrow \underline{X}^{\top} \beta+\beta^{(0)}-h^{-1}(1 / 2)>0$
- Linear Classifier: $\operatorname{sign}\left(\underline{X}^{\top} \beta+\beta^{(0)}-h^{-1}(1 / 2)\right)$


## Plugin Linear Classifier Estimation

- Classical choice for $h$ :

$$
\begin{aligned}
\lg \frac{G(t)}{1-h(t)}=t \longleftrightarrow h(t) & =\frac{e^{t}}{1+e^{t}} & \text { logit or logistic } \\
h(t) & =F_{\mathrm{N}}(t) & \text { probit } \\
h(t) & =1-e^{-e^{t}} & \text { log-log }
\end{aligned}
$$

- Choice of the best $\beta$ from the data.


## Maximum Likelihood Estimate

## Probabilistic Model

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


## Maximum Likelihood Approach

- Minimization of the negative log-likelihood:

$$
-\sum_{i=1}^{n} \log \left(\mathbb{P}\left(Y_{i} \mid \underline{X}_{i}\right)\right)=-\sum_{i=1}^{n}\left(\mathbf{1}_{Y_{i}=1} \log \left(h\left(\underline{X}_{i}^{\top} \beta\right)\right)+\mathbf{1}_{Y_{i}=-1} \log \left(1-h\left(\underline{X}_{i}^{\top} \beta\right)\right)\right)
$$

- Minimization possible if $h$ is regular...


## Maximum Likelihood Estimate

## KL Distance and negative log-likelihood

- Natural probalistic distance: Kullback-Leibler divergence

$$
\begin{aligned}
& \mathrm{KL}\left(\mathcal{B}(\mathbb{P}(Y=1 \mid \underline{X})), \mathcal{B}\left(h\left(\underline{X}^{\top} \beta\right)\right)\right. \\
& =\mathbb{E}_{\underline{X}}\left[\mathbb{P}(Y=1 \mid \underline{X}) \log \frac{\mathbb{P}(Y=1 \mid \underline{X})}{h\left(\underline{X}^{\top} \beta\right)}\right. \\
& \left.\quad+\mathbb{P}(Y=-1 \mid \underline{X}) \log \frac{1-\mathbb{P}(Y=1 \mid \underline{X})}{1-h\left(\underline{X}^{\top} \beta\right)}\right] \\
& =\mathbb{E}_{\underline{X}}\left[-\mathbb{P}(Y=1 \mid \underline{X}) \log \left(h\left(\underline{X}^{\top} \beta\right)\right)\right. \\
& \left.\quad-\mathbb{P}(Y=-1 \mid \underline{X}) \log \left(1-h\left(\underline{X}^{\top} \beta\right)\right)\right]+C_{\underline{X}, Y}
\end{aligned}
$$

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

$$
-\frac{1}{n} \sum_{i=1}^{n}\left(\mathbf{1}_{Y_{i}=1} \log \left(h\left(\underline{X}_{i}^{\top} \beta\right)\right)+\mathbf{1}_{Y_{i}=-1} \log \left(1-h\left(\underline{X}_{i}^{\top} \beta\right)\right)\right)
$$

Logistic Regression
Logistic Regression and Odd

- Logistic model: $h(t)=\frac{e^{t}}{1+e^{t}}$ (most natural choice... )
- The Bernoulli law $\mathcal{B}(h(t))$ satisfies then

$$
\frac{\mathbb{P}(Y=1)}{\mathbb{P}(Y=-1)}=e^{t} \Leftrightarrow \log \frac{\mathbb{P}(Y=1)}{\mathbb{P}(Y=-1)}=t
$$

- Interpretation in term of odd.
- Logistic model: linear model on the logarithm of the odd

$$
\log \frac{\mathbb{P}(Y=1 \mid \underline{X})}{\mathbb{P}(Y=-1 \mid \underline{X})}=\underline{X}^{\top} \beta
$$

## Associated Classifier

- Plugin strategy:

$$
f_{\beta}(\underline{X})= \begin{cases}1 & \text { if } \frac{e^{\underline{X}^{\top} \beta}}{1+e^{X^{\top} \beta}}>1 / 2 \Leftrightarrow \underline{X}^{\top} \beta>0 \\ -1 & \text { otherwise }\end{cases}
$$

## Likelihood Rewriting

- Negative log-likelihood:

$$
\begin{aligned}
& -\frac{1}{n} \sum_{i=1}^{n}\left(\mathbf{1}_{Y_{i}=1} \log \left(h\left(\underline{X}_{i}^{\top} \beta\right)\right)+\mathbf{1}_{Y_{i}=-1} \log \left(1-h\left(\underline{X}_{i}^{\top} \beta\right)\right)\right) \\
& =-\frac{1}{n} \sum_{i=1}^{n}\left(\mathbf{1}_{Y_{i}=1} \log \frac{e^{X_{i}^{\top} \beta}}{1+e^{\underline{X}_{i}^{\top} \beta}}+\mathbf{1}_{Y_{i}=-1} \log \frac{1}{1+e^{\underline{X}_{i}^{\top} \beta}}\right) \\
& =\frac{1}{n} \sum_{i=1}^{n} \log \left(1+e^{-Y_{i}\left(\underline{X}_{i}^{\top} \beta\right)}\right)
\end{aligned}
$$

- Convex and smooth function of $\beta$
- Easy optimization.

Logistic


## Transformed Representation

- From $\underline{X}$ to $\Phi(\underline{X})$ !
- New description of $\underline{X}$ leads to a different linear model:

$$
f_{\beta}(\underline{X})=\Phi(\underline{X})^{\top} \beta
$$

## Feature Design

- Art of choosing $\Phi$.
- Examples:
- Renormalization, (domain specific) transform
- Basis decomposition
- Interaction between different variables. . .

Example: Quadratic Logistic

$$
(A, B) \rightarrow\left(A, B, A, A,,_{1}^{2}, B^{2}\right)
$$

Quadratic Logistic




## Gaussian Linear Model

- Model: $Y \mid \underline{X} \sim \mathrm{~N}\left(\underline{X}^{\top} \beta, \sigma^{2}\right)$ plus independence
- Probably the most classical model of all time!
- Maximum Likelihood with explicit formulas for the two parameters.
- In regression, estimation of $\mathbb{E}[Y \mid \underline{X}]$ 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) $\left(v\left(\mathbb{E}_{\theta}[Y]\right)=\theta\right.$ 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: $f(y, \theta)=e^{y \theta-\ln \left(1+e^{\theta}\right)}(\theta=\ln p /(1-p))$
- Poisson: $f(y, \theta)=e^{\left(y \theta-e^{\theta}\right)+\ln (y!)}(\theta=\ln \lambda)$
- Linear Conditional model: $Y \mid \underline{X} \sim P_{\underline{x}^{\top} \beta} \ldots$
- Maximum likelihood fit of the parameters


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


## Non Parametric Conditional Estimation

- Two heuristics:
- $Y \mid \underline{X}$ (or $\mathbb{E}[Y \mid X]$ ) is almost constant (or simple) in a neighborhood of $\underline{X}$. (Kernel methods)
- $Y \mid \underline{X}$ (or $\mathbb{E}[Y \mid \underline{X}]$ ) 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 $Y \mid \underline{X}$ is locally constant or simple!


## Kernel

- Choose a kernel $K$ (think of a weighted neighborhood).
- For each $\underline{\tilde{X}}$, compute a simple localized estimate of $Y \mid \underline{X}$
- Use this local estimate to take the decision
- In regression, estimation of $\mathbb{E}[Y \mid \underline{X}]$ is sufficient.

Example: $k$ Nearest-Neighbors (with $k=3$ )


Example: $k$ Nearest-Neighbors (with $k=4$ )


- Neighborhood $\mathcal{V}_{\underline{x}}$ of $\underline{x}$ : $k$ learning samples closest from $\underline{x}$.
k-NN as local conditional density estimate

$$
\mathbb{P}(\widehat{Y=1} \mid \underline{X})=\frac{\sum_{\underline{X}_{i} \in \mathcal{V}_{\underline{X}}} \mathbf{1}_{\left\{Y_{i}=+1\right\}}}{\left|\mathcal{V}_{\underline{X}}\right|}
$$

- KNN Classifier:

$$
\widehat{f}_{K N N}(\underline{X})= \begin{cases}+1 & \text { if } \mathbb{P}(\widehat{Y=1} \underline{X}) \geq \mathbb{P}(\widehat{Y=-1} \mid \underline{X}) \\ -1 & \text { otherwise }\end{cases}
$$

- Lazy learning: all the computations have to be done at prediction time.
- Remark: You can also use your favorite kernel estimator...
$k-N N$ with $k=1$


$k-N N$ with $k=5$


$k-N N$ with $k=9$

k-NN with $\mathrm{k}=13$

$\mathrm{k}-\mathrm{NN}$ with $\mathrm{k}=17$


$k-N N$ with $\mathrm{k}=21$

k-NN with $\mathrm{k}=25$

$\mathrm{k}-\mathrm{NN}$ with $\mathrm{k}=29$

k-NN with $\mathrm{k}=33$

k-NN with $\mathrm{k}=37$

$\mathrm{k}-\mathrm{NN}$ with $\mathrm{k}=45$

k-NN with $\mathrm{k}=53$

k-NN with $\mathrm{k}=61$

k-NN with $\mathrm{k}=69$

k-NN with $\mathrm{k}=77$


$k-N N$ with $k=85$

$k-N N$ with $k=101$


k-NN with $\mathrm{k}=109$

k -NN with $\mathrm{k}=117$


k-NN with $\mathrm{k}=125$

$k-N N$ with $k=133$

$k-N N$ with $k=141$

$k-N N$ with $k=149$

k-NN with $\mathrm{k}=157$


k-NN with $\mathrm{k}=165$

k-NN with $\mathrm{k}=173$

k-NN with $\mathrm{k}=181$

k-NN with $\mathrm{k}=189$

k-NN with $\mathrm{k}=197$



## A naive idea

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

$$
\widehat{f}(\underline{X})=\frac{1}{\left|\left\{\underline{X}_{i} \in \mathcal{N}(\underline{X})\right\}\right|} \sum_{\underline{X}_{i} \in \mathcal{N}(\underline{X})} Y_{i}
$$

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

- Heuristic:
- If $\underline{X} \rightarrow \mathbb{E}[Y \mid \underline{X}]$ is regular then

$$
\mathbb{E}[Y \mid \underline{X}] \simeq \mathbb{E}\left[\mathbb{E}\left[Y \mid \underline{X}^{\prime}\right] \mid \underline{X}^{\prime} \in \mathcal{N}(\underline{X})\right]=\mathbb{E}\left[Y \mid \underline{X}^{\prime} \in \mathcal{N}(\underline{X})\right]
$$

- Replace an expectation by an empirical average:

$$
\mathbb{E}\left[Y \mid \underline{X}^{\prime} \in \mathcal{N}(\underline{X})\right] \simeq \frac{1}{\left|\left\{\underline{X}_{i} \in \mathcal{N}(\underline{X})\right\}\right|} \sum_{\underline{X}_{i} \in \mathcal{N}(\underline{X})} Y_{i}
$$

- Same idea than in classification where the proportion for class $k$ is estimated with the same formula by replacing $Y_{i}$ with $\mathbf{1}_{Y_{i}==k}$.


## Neighborhood and Size

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


## 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 $\mathbb{E}[Y \mid \underline{X}] \simeq \mathbb{E}\left[Y \mid \underline{X^{\prime}} \in \mathcal{N}(\underline{X})\right]$ is more accurate (small bias).


## Weighted Local Averaging

## Weighted Local Average

- Replace the neighborhood $\mathcal{N}(\underline{X})$ by a decaying window function $w\left(\underline{X}, \underline{X}^{\prime}\right)$.
- $\mathbb{E}[Y \mid \underline{X}]$ can be approximated by a weighted local average:

$$
\hat{f}(\underline{X})=\frac{\sum_{i} w\left(\underline{X}, \underline{X}_{i}^{\prime}\right) Y_{i}}{\sum_{i} w\left(\underline{X}, \underline{X}_{i}^{\prime}\right)} .
$$

## Kernel

- Most classical choice: $w\left(\underline{X}, \underline{X}^{\prime}\right)=K\left(\frac{\underline{X}-\underline{X}^{\prime}}{h}\right)$ where $h$ the bandwidth is a scale parameter.
- Examples:
- Box kernel: $K(t)=\mathbf{1}_{\|t\| \leq 1}$ (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.


## A Density Estimation Point of View?

## Nadaraya-Watson Heuristic

- Provided all the densities exist

$$
\mathbb{E}[Y \mid \underline{X}]=\frac{\int Y p(\underline{X}, Y) d Y}{\int p(Y, \underline{X}) d Y}=\frac{\int Y p(\underline{X}, Y) d Y}{p(\underline{X})}
$$

- Replace the unknown densities by their estimates:

$$
\begin{aligned}
\hat{p}(\underline{X}) & =\frac{1}{n} \sum_{i=1}^{n} K\left(\underline{X}-\underline{X}_{i}\right) \\
\hat{p}(\underline{X}, Y) & =\frac{1}{n} \sum_{i=1}^{n} K\left(\underline{X}-\underline{X}_{i}\right) K^{\prime}\left(Y-Y_{i}\right)
\end{aligned}
$$

- Now if $K^{\prime}$ is a kernel such that $\int Y K^{\prime}(Y) d Y=0$ then

$$
\int Y \widehat{p}(\underline{X}, Y) d Y=\frac{1}{n} \sum_{i=1}^{n} K\left(\underline{X}-\underline{X}_{i}\right) Y_{i}
$$

## A Density Estimation Point of View?

## Nadaraya-Watson

- Resulting estimator of $\mathbb{E}[Y \mid \underline{X}]$

$$
\widehat{f}(\underline{X})=\frac{\sum_{i=1}^{n} Y_{i} K_{h}\left(\underline{X}-\underline{X}_{i}\right)}{\sum_{i=1}^{n} K_{h}\left(\underline{X}-\underline{X}_{i}\right)}
$$

- 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!


## Another Point of View on Kernel

- Nadaraya-Watson estimator:

$$
\widehat{f}(\underline{X})=\frac{\sum_{i=1}^{n} Y_{i} K_{h}\left(\underline{X}-\underline{X}_{i}\right)}{\sum_{i=1}^{n} K_{h}\left(\underline{X}-\underline{X}_{i}\right)}
$$

- Can be view as a minimizer of

$$
\sum_{i=1}^{n}\left|Y_{i}-\beta\right|^{2} K_{h}\left(\underline{X}-\underline{X}_{i}\right)
$$

- Local regression of order 0 .


## Local Linear Model

- Estimate $\mathbb{E}[Y \mid \underline{X}]$ by $\widehat{f}(\underline{X})=\phi(\underline{X})^{\top} \widehat{\beta}(\underline{X})$ where $\phi$ is any function of $\underline{X}$ and $\widehat{\beta}(\underline{X})$ is the minimizer of

$$
\sum_{i=1}^{n}\left|Y_{i}-\phi\left(\underline{X}_{i}\right)^{\top} \beta\right|^{2} K_{h}\left(\underline{X}-\underline{X}_{i}\right)
$$

- Very similar to a piecewise modeling approach.


## LOESS: LOcal polynomial regrESSion

## 1D Nonparametric Regression

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

$$
\sum_{i=1}^{n}\left|Y_{i}-\sum_{j=0}^{d} \beta^{(j)} \underline{X}_{i}^{j}\right|^{2} K_{h}\left(\underline{X}-\underline{X}_{i}\right) .
$$

- Most classical kernel used: Tricubic kernel

$$
K(t)=\max \left(1-|t|^{3}, 0\right)^{3}
$$

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


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- Non Parametric Conditional Density Modeling


## - Generative Modeling

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## Fully Generative Modeling

- Idea: If one knows the law of $(\underline{X}, Y)$ everything is easy!


## Bayes formula

- With a slight abuse of notation,

$$
\begin{aligned}
\mathbb{P}(Y \mid \underline{X}) & =\frac{\mathbb{P}((\underline{X}, Y))}{\mathbb{P}(\underline{X})} \\
& =\frac{\mathbb{P}(\underline{X} \mid Y) \mathbb{P}(Y)}{\mathbb{P}(\underline{X})}
\end{aligned}
$$

$$
\begin{aligned}
&(x, y) \\
&=(1 y \mid x) P(x) \\
&=P(x(y) P(y)
\end{aligned}
$$

- Generative Modeling:
- Propose a model for $(\underline{X}, Y)$ (or equivalently $\underline{X} \mid Y$ 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 $(\underline{X}, Y)$ rather than only $Y \mid \underline{X}$ !
- Great flexibility in the model design but may lead to complex computation.


## Fully Generative Modeling

- Simpler setting in classification!


## Bayes formula

$$
\mathbb{P}(Y=k \mid \underline{X})=\frac{\mathbb{P}(\underline{X} \mid Y=k) \mathbb{P}(Y=k)}{\mathbb{P}(X)}
$$

- Binary Bayes classifier (the best solution)

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

- Heuristic: Estimate those quantities and plug the estimations.
- By using different models/estimators for $\mathbb{P}(\underline{X} \mid Y)$, we get different classifiers.
- Rk: No need to renormalize by $\mathbb{P}(\underline{X})$ to take the decision!

Discriminant Analysis

## Discriminant Analysis (Gaussian model)

- The densities are modeled as multivariate normal, i.e.,

$$
\mathbb{P}(\underline{X} \mid Y=k) \sim \mathrm{N}_{\mu_{k}, \Sigma_{k}}
$$

- Discriminant functions: $\boldsymbol{g}_{\boldsymbol{k}}(\underline{X})=\ln (\mathbb{P}(\underline{X} \mid Y=k))+\ln (\mathbb{P}(Y=k))$

$$
\begin{aligned}
g_{k}(\underline{X})=- & \frac{1}{2}\left(\underline{X}-\mu_{k}\right)^{\top} \Sigma_{k}^{-1}\left(\underline{X}-\mu_{k}\right) \\
& -\frac{d}{2} \ln (2 \pi)-\frac{1}{2} \ln \left(\left|\Sigma_{k}\right|\right)+\ln (\mathbb{P}(Y=k))
\end{aligned}
$$

- 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!



## Quadratic 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



## Quadratic Discriminant Analysis

- The probability densities are Gaussian
- The effect of any decision rule is to divide the feature space into some decision regions $\mathcal{R}_{1}, \mathcal{R}_{2}, \ldots, \mathcal{R}_{c}$
- The regions are separated by decision boundaries


## Estimation

In practice, we will need to estimate $\mu_{k}, \Sigma_{k}$ and $\mathbb{P}_{k}:=\mathbb{P}(Y=k)$

- The estimate proportion $\mathbb{P}(\widehat{Y=k})=\frac{n_{k}}{n}=\frac{1}{n} \sum_{i=1}^{n} \mathbf{1}_{\left\{Y_{i}=k\right\}}$
- Maximum likelihood estimate of $\widehat{\mu_{k}}$ and $\widehat{\Sigma_{k}}$ (explicit formulas)
- DA classifier

$$
\widehat{f}_{G}(\underline{X})= \begin{cases}+1 & \text { if } \widehat{g}_{+1}(\underline{X}) \geq \widehat{g}_{-1}(\underline{X}) \\ -1 & \text { otherwise }\end{cases}
$$

- 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

- $\Sigma_{\omega_{1}}=\Sigma_{\omega_{2}}=\Sigma$
- The decision boundaries are linear hyperplanes



## Quadratic Discriminant Analysis

- $\Sigma_{\omega_{1}} \neq \Sigma_{\omega_{2}}$
- Arbitrary Gaussian distributions lead to Bayes decision boundaries that are general quadratics.

Linear Discrimant Analysis


Quadratic Discrimant Analysis

Decision region


Decision boundary


## Naive Bayes

- Classical algorithm using a crude modeling for $\mathbb{P}(\underline{X} \mid Y)$ :
- Feature independence assumption:

$$
\mathbb{P}(\underline{X} \mid Y)=\prod_{l=1}^{d} \mathbb{P}\left(\underline{X}^{(\prime)} \mid Y\right)
$$

- 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!

Naive Bayes with Gaussian model


Naive Bayes with Density Estimation


Naive Bayes with kernel density estimates


## 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

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

Canonical approach: $\widehat{f}_{\mathcal{S}}=\operatorname{argmin}_{f \in \mathcal{S}} \frac{1}{n} \sum_{i=1}^{n} \ell\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)$

## Problems

- How to choose $\mathcal{S}$ ?
- How to compute the minimization?


## A Probabilistic Point of View

Solution: For $\underline{X}$, estimate $Y \mid \underline{X}$ 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 $f_{\theta}(\underline{X})=\underline{X}^{\top} \beta+\beta^{(0)}$ with $\theta=\left(\beta, \beta^{(0)}\right)$.
- Find $\hat{\theta}=\arg \min \frac{1}{n} \sum_{i=1}^{n} \log \left(1+e^{-Y_{i} f_{\theta}\left(X_{i}\right)}\right)+\lambda\|\beta\|_{1}$
- Classify using $\operatorname{sign}\left(f_{\hat{\theta}}\right)$


## Deep Learning

- Let $f_{\theta}(\underline{X})$ 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 $-\frac{1}{n} \sum_{i=1}^{n} \log \left(f_{\theta}\left(\underline{X}_{i}\right)^{\left(Y_{i}\right)}\right)$
- Classify using $\operatorname{sign}\left(f_{\hat{\theta}}\right)$


## Support Vector Machine

- Let $f_{\theta}(\underline{X})=\underline{X}^{\top} \beta+\beta^{(0)}$ with $\theta=\left(\beta, \beta^{(0)}\right)$.
- Find $\hat{\theta}=\arg \min \frac{1}{n} \sum_{i=1}^{n} \max \left(1-Y_{i} f_{\theta}\left(\underline{X}_{i}\right), 0\right)+\lambda\|\beta\|_{2}^{2}$
- Classify using $\operatorname{sign}\left(f_{\hat{\theta}}\right)$
- Those three methods rely on a similar heuristic: the optimization point of view!
- The best solution $f^{\star}$ is the one minimizing

$$
f^{\star}=\arg \min R(f)=\arg \min \mathbb{E}[\ell(Y, f(\underline{X}))]
$$

## Empirical Risk Minimization

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

$$
\widehat{f}=f_{\widehat{\theta}}=\underset{f_{\theta}, \theta \in \Theta}{\operatorname{argmin}} \frac{1}{n} \sum_{i=1}^{n} \ell\left(Y_{i}, f_{\theta}\left(\underline{X}_{i}\right)\right)
$$

- Intractable for the $\ell^{0 / 1}$ loss!


## Convexification Strategy

## Risk Convexification

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

$$
\tilde{f}=f_{\widehat{\theta}}=\underset{f_{\theta}, \theta \in \Theta}{\operatorname{argmin}} \frac{1}{n} \sum_{i=1}^{n} \bar{\ell}\left(Y_{i}, f_{\theta}\left(\underline{X}_{i}\right)\right)
$$

- Use $\widehat{f}=\operatorname{sign}(\tilde{f})$
- Much easier optimization.


## Instantiation

- Logistic (Revisited)
- Support Vector Machine
- (Deep) Neural Network
- Boosting


## Classification Loss and Convexification



## Convexification

- Replace the loss $\ell^{0 / 1}(Y, f(\underline{X}))$ by

$$
\bar{\ell}(Y, f(\underline{X}))=I(Y f(\underline{X}))
$$

with / a convex function.

- Further mild assumption: $I$ is decreasing, differentiable at 0 and $I^{\prime}(0)<0$.


## Classification Loss and Convexification



## Classical convexification

- Logistic loss: $\bar{\ell}(Y, f(\underline{X}))=\log _{2}\left(1+e^{-Y f(\underline{X})}\right)($ Logistic $/ N N)$
- Hinge loss: $\bar{\ell}(Y, f(\underline{X}))=(1-Y f(\underline{X}))_{+}($SVM $)$
- Exponential loss: $\bar{\ell}(Y, f(\underline{X}))=e^{-Y f(\underline{X})}$ (Boosting...)


## Properties

## The Target is the Bayes Classifier

- The minimizer of

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

is the Bayes classifier $f^{\star}=\operatorname{sign}(2 \eta(\underline{X})-1)$

## Control of the Excess Risk

- It exists a convex function $\Psi$ such that

$$
\begin{gathered}
\Psi\left(\mathbb { E } \left[\ell^{0 / 1}(Y, \operatorname{sign}(f(\underline{X}))]-\mathbb{E}\left[\ell^{0 / 1}\left(Y, f^{\star}(\underline{X})\right]\right)\right.\right. \\
\leq \mathbb{E}\left[\bar{\ell}(Y, f(\underline{X})]-\mathbb{E}\left[\bar{\ell}\left(Y, f^{\star}(\underline{X})\right)\right]\right.
\end{gathered}
$$

- Theoretical guarantee!
- Ideal solution:

$$
\widehat{f}=\underset{f \in \mathcal{S}}{\operatorname{argmin}} \frac{1}{n} \sum_{i=1}^{n} \ell^{0 / 1}\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)
$$

## Logistic regression

- Use $f(\underline{X})=\underline{X}^{\top} \beta+\beta^{(0)}$.
- Use the logistic loss $\bar{\ell}(y, f)=\log _{2}\left(1+e^{-y f}\right)$, i.e. the negative log-likelihood.
- Different vision than the statistician but same algorithm!


## Logistic




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## 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 $h\left(\underline{x}^{\top} \beta\right)$.


## 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 $\|\beta\|_{0}=\sum_{i=1}^{d} \mathbf{1}_{\beta^{(i)} \neq 0}<C$
- Norm: Impose that $\|\beta\|_{p}<C$ with $1 \leq p($ Often $p=2$ or $p=1)$

Norms and Sparsity


## Sparsity

- $\beta$ is sparse if its number of non-zero coefficients $\left(\ell_{0}\right)$ 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

$$
\underset{\beta \in \mathbb{R}^{d},\|\beta\|_{p} \leq C}{\operatorname{argmin}} \frac{1}{n} \sum_{i=1}^{n} \bar{\ell}\left(Y_{i}, h\left(\underline{x}_{i}^{\top} \beta\right)\right)
$$

## Lagrangian Reformulation

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

$$
\underset{\beta \in \mathbb{R}^{d}}{\operatorname{argmin}} \frac{1}{n} \sum_{i=1}^{n} \bar{\ell}\left(Y_{i}, h\left(\underline{x}_{i}^{\top} \beta\right)\right)+\lambda\|\beta\|_{p}^{p^{\prime}}
$$

with $p^{\prime}=p$ except if $p=0$ where $p^{\prime}=1$.

- Easier calibration. . . but no explicit model $\mathcal{S}$.
- $\mathbf{R k}:\|\beta\|_{p}$ is not scaling invariant if $p \neq 0 \ldots$
- Initial rescaling issue.


## Penalization

## Penalized Linear Model

- Minimization of

$$
\underset{\beta \in \mathbb{R}^{d}}{\operatorname{argmin}} \frac{1}{n} \sum_{i=1}^{n} \bar{\ell}\left(Y_{i}, h\left(\underline{x}_{i}^{\top} \beta\right)\right)+\operatorname{pen}(\beta)
$$

where $\operatorname{pen}(\beta)$ is a (sparsity promoting) penalty

- Variable selection if $\beta$ is sparse.


## Classical Penalties

- AIC: $\operatorname{pen}(\beta)=\lambda\|\beta\|_{0}$ (non-convex / sparsity)
- Ridge: $\operatorname{pen}(\beta)=\lambda\|\beta\|_{2}^{2}$ (convex / no sparsity)
- Lasso: $\operatorname{pen}(\beta)=\lambda\|\beta\|_{1}$ (convex / sparsity)
- Elastic net: $\operatorname{pen}(\beta)=\lambda_{1}\|\beta\|_{1}+\lambda_{2}\|\beta\|_{2}^{2}$ (convex / sparsity)
- Easy optimization if pen (and the loss) is convex...
- Need to specify $\lambda$ to define a ML method!


## Classical Examples

- Penalized Least Squares
- Penalized Logistic Regression
- Penalized Maximum Likelihood
- SVM
- Tree pruning
- Sometimes used even if the parameterization is not linear. .


## Practical Selection Methodology

- Choose a penalty family pen ${ }_{\lambda}$.
- Compute a CV risk for the penalty pen ${ }_{\lambda}$ for all $\lambda \in \Lambda$.
- Determine $\widehat{\lambda}$ the $\lambda$ minimizing the CV risk.
- Compute the final model with the penalty pen $\hat{\lambda}$.
- CV allows to select a ML method, penalized estimation with a penalty pen $\widehat{\lambda}$, 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...)


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## Perceptron



## Perceptron (Rosenblatt 1957)

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


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# 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.


## Logistic unit

- Structure:
- Mix inputs with a weighted sum,
- Apply the logistic function $\sigma(t)=e^{t} /\left(1+e^{t}\right)$,
- Threshold at $1 / 2$ to make a decision!
- Logistic weights learned by minimizing the -log-likelihood.
- Equivalent to linear regression when using a linear activation function!



## 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!

Neural Network


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!

H2O NN




## 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!


## Convolutional Network

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Fig. 2. Architecture of LeNet-5, a Convolutional Neural Network, here for digits recognition. Each plane is a feature map, i.e. a set of units whose weights are constrained to be identical.

## 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 60000 examples!
- Tremendous improvement.
- Representation learned through the task.



## Alexnet - A. Krizhevsky, I. Sutskever, G. Hinton (2012)

- Bigger and deeper layers and thus much more parameters.
- Clever intialization 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 Progess!


## Outline

A Probabilistic Point of View(1)

Introduction Machine Learning

- Motivation
(2)

A Practical View

- Method or Models
- Interpretability
- Metric Choice
(3)

A Better Point of View

- The Example of Univariate Linear Regression
- Supervised LearningRisk Estimation and Method Choice - Cross Validation
- Cross Validation and Test
- Cross Validation and Weights
- Auto ML
- Parametric Conditional Density Modeling
- Non Parametric Conditional Density Modeling
- Generative Modeling
(6) Optimization Point of View
- Penalization
- (Deep) Neural Networks
- SVM
- Tree Based Methods
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(D) Empirical Risk Minimization
- Empirical Risk Minimization
- ERM and PAC Bayesian Analysis
- Hoeffding and Finite Class
- McDiarmid and Rademacher Complexity
- VC Dimension
- Structural Risk Minimization
(8) References

$$
\begin{aligned}
f_{\theta}(\underline{X}) & =\underline{X}^{\top} \beta+\beta^{(0)} \quad \text { with } \quad \theta=\left(\beta, \beta^{(0)}\right) \\
\hat{\theta} & =\arg \min \frac{1}{n} \sum_{i=1}^{n} \max \left(1-Y_{i} f_{\theta}\left(\underline{X}_{i}\right), 0\right)+\lambda\|\beta\|_{2}^{2}
\end{aligned}
$$

## Support Vector Machine

- Convexification of the 0/1-loss with the hinge loss:

$$
\mathbf{1}_{Y_{i} f_{\theta}\left(\underline{X}_{i}\right)<0} \leq \max \left(1-Y_{i} f_{\theta}\left(\underline{X}_{i}\right), 0\right)
$$

- Penalization by the quadratic norm (Ridge/Tikhonov).
- Solution can be approximated by gradient descent algorithms.
- Revisit of the original point of view.
- Original point of view leads to a different optimization algorithm and to some extensions.

- Linear classifier: $\operatorname{sign}\left(\underline{X}^{\top} \beta+\beta^{(0)}\right)$
- Separable case: $\exists\left(\beta, \beta^{(0)}\right), \forall i, Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right)>0$

How to choose $\left(\beta, \beta^{(0)}\right)$ so that the separation is maximal?

- Strict separation: $\exists\left(\beta, \beta^{(0)}\right), \forall i, Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right) \geq 1$
- Distance between $\underline{X}^{\top} \beta+\beta^{(0)}=1$ and $\underline{X}^{\top} \beta+\beta^{(0)}=-1$ :

$$
\frac{2}{\|\beta\|}
$$

- Maximizing this distance is equivalent to minimizing $\frac{1}{2}\|\beta\|^{2}$.



## Separable SVM

- Constrained optimization formulation:

$$
\min \frac{1}{2}\|\beta\|^{2} \quad \text { with } \quad \forall i, Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right) \geq 1
$$

- Quadratic Programming setting.
- Efficient solver available...

Non Separable Case


- What about the non separable case?


## SVM relaxation

- Relax the assumptions

$$
\forall i, Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right) \geq 1 \text { to } \forall i, Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right) \geq 1-s_{i}
$$

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:

$$
\min \frac{1}{2}\|\beta\|^{2}+C \sum_{i=1}^{n} s_{i} \quad \text { with } \quad\left\{\begin{array}{l}
\forall i, Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right) \geq 1-s_{i} \\
\forall i, s_{i} \geq 0
\end{array}\right.
$$

- Hinge Loss reformulation:

$$
\min \frac{1}{2}\|\beta\|^{2}+C \sum_{i=1}^{n} \underbrace{\max \left(0,1-Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right)\right)}_{\text {Hinge Loss }}
$$

- Constrained convex optimization algorithms vs gradient descent algorithms.
- Convex relaxation:

$$
\begin{aligned}
& \operatorname{argmin} \frac{1}{2}\|\beta\|^{2}+C \sum_{i=1}^{n} \max \left(1-Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right), 0\right) \\
& =\operatorname{argmin} \frac{1}{n} \sum_{i=1}^{n} \max \left(1-Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right), 0\right)+\frac{1}{C n} \frac{1}{2}\|\beta\|^{2}
\end{aligned}
$$

- Prop: $\ell^{0 / 1}\left(Y_{i}, \operatorname{sign}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right)\right) \leq \max \left(1-Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right), 0\right)$


## Penalized convex relaxation (Tikhonov!)

$$
\begin{aligned}
& \frac{1}{n} \sum_{i=1}^{n} \ell^{0 / 1}\left(Y_{i}, \operatorname{sign}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right)\right)+\frac{1}{C n} \frac{1}{2}\|\beta\|^{2} \\
& \quad \leq \frac{1}{n} \sum_{i=1}^{n} \max \left(1-Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right), 0\right)+\frac{1}{C n} \frac{1}{2}\|\beta\|^{2}
\end{aligned}
$$

Support Vector Machine
Decision region


## Constrained Minimization

## Constrained Minimization

- Goal:

$$
\begin{aligned}
& \min _{x} f(x) \\
& \text { with } \begin{cases}h_{j}(x)=0, & j=1, \ldots p \\
g_{i}(x) \leq 0, & i=1, \ldots q\end{cases}
\end{aligned}
$$

- or rather with 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


## Constrained Minimization

- Goal:

$$
p^{\star}=\min _{x} f(x) \quad \text { with } \quad \begin{cases}h_{j}(x)=0, & j=1, \ldots p \\ g_{i}(x) \leq 0, & i=1, \ldots q\end{cases}
$$

## 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 $\mu \in\left(\mathbb{R}^{+}\right)^{q}$.

- The $\lambda_{j}$ and $\mu_{i}$ are called the dual (or Lagrange) variables.
- Prop:

$$
\begin{aligned}
\max _{\lambda \in \mathbb{R}^{p}, \mu \in\left(\mathbb{R}^{+}\right)^{q}} \mathcal{L}(x, \lambda, \mu) & = \begin{cases}f(x) & \text { if } x \text { is feasible } \\
+\infty & \text { otherwise }\end{cases} \\
\min _{x} \max _{\lambda \in \mathbb{R}^{p}, \mu \in\left(\mathbb{R}^{+}\right)^{q}} \mathcal{L}(x, \lambda, \mu) & =p^{\star}
\end{aligned}
$$

## 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 $\mu \in\left(\mathbb{R}^{+}\right)^{q}$.

## Lagrangian Dual

- Lagrangian dual function:

$$
Q(\lambda, \mu)=\min _{x} \mathcal{L}(x, \lambda, \mu)
$$

- Prop:

$$
\begin{aligned}
& Q(\lambda, \mu) \leq f(x) \text {, for all feasible } x \\
& \max _{\lambda \in \mathbb{R}^{P}, \mu \in\left(\mathbb{R}^{+}\right)^{q}} Q(\lambda, \mu) \leq \min _{x \text { feasible }} f(x)
\end{aligned}
$$

## Duality

## Primal

- Primal:

$$
p^{\star}=\min _{x \in \mathcal{X}} f(x) \text { with } \begin{cases}h_{j}(x)=0, & j=1, \ldots p \\ g_{i}(x) \leq 0, & i=1, \ldots q\end{cases}
$$

## Dual

- Dual:

$$
q^{\star}=\max _{\lambda \in \mathbb{R}^{p}, \mu \in\left(\mathbb{R}^{+}\right)^{q}} Q(\lambda, \mu)=\max _{\lambda \in \mathbb{R}^{p}, \mu \in\left(\mathbb{R}^{+}\right)^{q}} \min _{x} \mathcal{L}(x, \lambda, \mu)
$$

## Duality

- Always weak duality:

$$
\begin{aligned}
q^{\star} & \leq p^{\star} \\
\max _{\lambda \in \mathbb{R}^{p}, \mu \in\left(\mathbb{R}^{+}\right)^{q}} \min _{x} \mathcal{L}(x, \lambda, \mu) & \leq \min _{x} \max _{\lambda \in \mathbb{R}^{p}, \mu \in\left(\mathbb{R}^{+}\right)^{q}} \mathcal{L}(x, \lambda, \mu)
\end{aligned}
$$

- Not always strong duality $q^{\star}=p^{\star}$.


## Strong Duality

## Strong Duality

- Strong duality:

$$
\begin{aligned}
q^{\star} & =p^{\star} \\
\max _{\lambda \in \mathbb{R}^{p}, \mu \in\left(\mathbb{R}^{+}\right)^{q}} \min _{x} \mathcal{L}(x, \lambda, \mu) & =\min _{x} \max _{\lambda \in \mathbb{R}^{p},{ }_{\mu \in\left(\mathbb{R}^{+}\right)^{q}} \mathcal{L}(x, \lambda, \mu)}
\end{aligned}
$$

- 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 Condition

- Stationarity:

$$
\nabla_{x} \mathcal{L}\left(x^{\star}, \lambda, \mu\right)=\nabla f\left(x^{\star}\right)+\sum_{j} \lambda_{j} \nabla h_{j}\left(x^{\star}\right)+\sum_{i} \mu_{i} \nabla g_{i}\left(x^{\star}\right)=0
$$

- Primal admissibility:

$$
h_{j}\left(x^{\star}\right)=0 \quad \text { and } \quad g_{i}\left(x^{\star}\right) \leq 0
$$

- Dual admissibility:

$$
\mu_{i} \geq 0
$$

- Complementary slackness:

$$
\mu_{i} g_{i}\left(x^{\star}\right)=0
$$

## KKT Theorem

- If $f$ convex, $h_{j}$ affine and $g_{i}$ convex, all are differentiable and strong duality holds then $x^{\star}$ is a solution of the primal problem if and only if the KKT condition holds


## SVM

- Constrained optimization formulation:

$$
\min \frac{1}{2}\|\beta\|^{2}+C \sum_{i=1}^{n} s_{i} \text { with }\left\{\begin{array}{l}
\forall i, Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right) \geq 1-s_{i} \\
\forall i, s_{i} \geq 0
\end{array}\right.
$$

## SVM Lagrangian

- Lagrangian:

$$
\begin{aligned}
\mathcal{L}\left(\beta, \beta^{(0)}, s, \alpha, \mu\right) & =\frac{1}{2}\|\beta\|^{2}+C \sum_{i=1}^{n} s_{i} \\
& +\sum_{i} \alpha_{i}\left(1-s_{i}-Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right)\right)-\sum_{i} \mu_{i} s_{i}
\end{aligned}
$$

## SVM and KKT

## KKT Optimality Conditions

- Stationarity:

$$
\begin{aligned}
\nabla_{\beta} \mathcal{L}\left(\beta, \beta^{(0)}, s, \alpha, \mu\right) & =\beta-\sum_{i} \alpha_{i} Y_{i} \underline{X}_{i}=0 \\
\nabla_{\beta^{(0)}} \mathcal{L}\left(\beta, \beta^{(0)}, s, \alpha, \mu\right) & =-\sum_{i} \alpha_{i}=0 \\
\nabla_{s_{i}} \mathcal{L}\left(\beta, \beta^{(0)}, s, \alpha, \mu\right) & =C-\alpha_{i}-\mu_{i}=0
\end{aligned}
$$

- Primal and dual admissibility:

$$
\left(1-s_{i}-Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right)\right) \leq 0, \quad s_{i} \geq 0, \quad \alpha_{i} \geq 0, \quad \text { and } \mu_{i} \geq 0
$$

- Complementary slackness:

$$
\alpha_{i}\left(1-s_{i}-Y_{i}\left(\underline{X}_{i}^{\top} \beta+\beta^{(0)}\right)\right)=0 \quad \text { and } \quad \mu_{i} s_{i}=0
$$

## Consequence

- $\beta^{\star}=\sum_{i} \alpha_{i} Y_{i} \underline{X}_{i}$ and $0 \leq \alpha_{i} \leq C$.
- If $\alpha_{i} \neq 0, \underline{X}_{i}$ is called a support vector and either
- $s_{i}=0$ and $Y_{i}\left(\underline{X}_{i}^{\top} \beta^{\star}+\beta^{(0) *}\right)=1$ (margin hyperplane),
- or $\alpha_{i}=C$ (outliers).
- $\beta^{(0) *}=Y_{i}-\underline{X}_{i}^{\top} \beta^{\star}$ for any support vector with $0<\alpha_{i}<C$.


## SVM Dual

## SVM Lagrangian Dual

- Lagrangian Dual:

$$
Q(\alpha, \mu)=\min _{\beta, \beta^{(0)}, s} \mathcal{L}\left(\beta, \beta^{(0)}, s, \alpha, \mu\right)
$$

- 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 $\forall i, \alpha_{i}+\mu_{i}=C$,

$$
Q(\alpha, \mu)=\sum_{i} \alpha_{i}-\frac{1}{2} \sum_{i, j} \alpha_{i} \alpha_{j} Y_{i} Y_{j} \underline{X}_{i}^{\top} \underline{X}_{j}
$$

## SVM Dual problem

- Dual problem is a Quadratic Programming problem:

$$
\max _{\alpha \geq 0, \mu \geq 0} Q(\alpha, \mu) \Leftrightarrow \max _{0 \leq \alpha \leq C} \sum_{i} \alpha_{i}-\frac{1}{2} \sum_{i, j} \alpha_{i} \alpha_{j} Y_{i} Y_{j} \underline{X}_{i}^{\top} \underline{X}_{j}
$$

- Involves the $\underline{X}_{i}$ 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

$$
\sum_{i=1}^{n} \bar{\ell}\left(Y_{i}, \underline{X}_{i}^{\top} \beta+\beta^{(0)}\right)+\Phi\left(\|\beta\|_{2}\right)
$$

is a linear combination of the input points $\beta^{\star}=\sum_{i=1}^{n} \alpha_{i}^{\prime} \underline{X}_{i}$.

- Minimization problem in $\alpha^{\prime}$ :

$$
\sum_{i=1}^{n} \bar{\ell}\left(Y_{i}, \sum_{j} \alpha_{j}^{\prime} \underline{X}_{i}^{\top} \underline{X}_{j}+\beta^{(0)}\right)+\Phi\left(\|\beta\|_{2}\right)
$$

involving only the scalar product of the data.

- Optimal predictor requires only to compute scalar products.

$$
\hat{f}^{\star}(\underline{X})=\underline{X}^{\top} \beta^{\star}+\beta^{(0), *}=\sum_{i} \alpha_{i}^{\prime} \underline{X}_{i}^{\top} \underline{X}
$$

- Transform a problem in dimension $\operatorname{dim}(\mathcal{X})$ 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 $\underline{X}$.
- Example: add monomials $\left(\underline{X}^{(j)}\right)^{2}, \underline{X}^{(j)} \underline{X}^{\left(j^{\prime}\right)} \ldots$
- Adding feature increases the dimension.


## Feature Map

- Application $\phi: \mathcal{X} \rightarrow \mathbb{H}$ with $\mathbb{H}$ an Hilbert space.
- Linear decision boundary in $\mathbb{H}: \phi(\underline{X})^{\top} \beta+\beta^{(0)}=0$ is not an hyperplane anymore in $\mathcal{X}$.
- Heuristic: Increasing dimension allows to make data almost linearly separable.

Polynomial Mapping


| $\substack{(1,1,+\sqrt{2},-\sqrt{2},-\sqrt{2}, 1)} \stackrel{\sqrt{2} x_{1} x_{2}}{\substack{(1,1,+\sqrt{2},+\sqrt{2},+\sqrt{2}, 1)}} \xrightarrow{\bullet} \sqrt{2} x_{1}$ |
| :---: |

## Polynomial Mapping of order 2

- $\phi: \mathbb{R}^{2} \rightarrow \mathbb{R}^{6}$

$$
\phi(\underline{X})=\left(\left(\underline{X}^{(1)}\right)^{2},\left(\underline{X}^{(2)}\right)^{2}, \sqrt{2} \underline{X}^{(1)} \underline{X}^{(2)}, \sqrt{2} \underline{X}^{(1)}, \sqrt{2} \underline{X}^{(2)}, 1\right)
$$

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


## Polynomial Mapping and Scalar Product

- Prop:

$$
\phi(\underline{X})^{\top} \phi\left(\underline{X}^{\prime}\right)=\left(1+\underline{X}^{\top} \underline{X}^{\prime}\right)^{2}
$$

## SVM Primal and Dual

## Primal, Lagrandian and Dual

- Primal:

$$
\min \|\beta\|^{2}+C \sum_{i=1}^{n} s_{i} \quad \text { with } \quad\left\{\begin{array}{l}
\forall i, Y_{i}\left(\phi\left(X_{i}\right)^{\top} \beta+\beta^{(0)}\right) \geq 1-s_{i} \\
\forall i, s_{i} \geq 0
\end{array}\right.
$$

- Lagrangian:

$$
\begin{aligned}
\mathcal{L}\left(\beta, \beta^{(0)}, s, \alpha, \mu\right) & =\frac{1}{2}\|\beta\|^{2}+C \sum_{i=1}^{n} s_{i} \\
& +\sum_{i} \alpha_{i}\left(1-s_{i}-Y_{i}\left(\phi\left(\underline{X}_{i}\right)^{\top} \beta+\beta^{(0)}\right)\right)-\sum_{i} \mu_{i} s_{i}
\end{aligned}
$$

- Dual:

$$
\max _{\alpha \geq 0, \mu \geq 0} Q(\alpha, \mu) \Leftrightarrow \max _{0 \leq \alpha \leq C} \sum_{i} \alpha_{i}-\frac{1}{2} \sum_{i, j} \alpha_{i} \alpha_{j} Y_{i} Y_{j} \phi\left(\underline{X}_{i}\right)^{\top} \phi\left(\underline{X}_{j}\right)
$$

- Optimal $\phi(\underline{X})^{\top} \beta^{\star}+\beta^{(0), *}=\sum_{i} \alpha_{i} Y_{i} \phi(\underline{X})^{\top} \phi\left(\underline{X}_{i}\right)$
- Only need to know to compute $\phi(\underline{X})^{\top} \phi\left(\underline{X}^{\prime}\right)$ to obtain the solution.


## From Map to Kernel

- Many algorithms (e.g. SVM) require only to be able to compute the scalar product $\phi(\underline{X})^{\top} \phi\left(\underline{X}^{\prime}\right)$.


## Kernel

- Any application

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

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

## Kernel Trick

- Computing directly the kernel $k\left(\underline{X}, \underline{X}^{\prime}\right)=\phi(\underline{X})^{\top} \phi\left(\underline{X}^{\prime}\right)$ may be easier than computing $\phi(\underline{X}), \phi\left(\underline{X}^{\prime}\right)$ and then the scalar product.
- Here $k$ is defined from $\phi$.
- Under some assumption on $k, \phi$ can be implicitly defined from $k$ !


## Positive Definite Symmetric Kernels

- A kernel $k$ is PDS if and only if
- $k$ is symmetric, i.e.

$$
k\left(\underline{X}, \underline{X}^{\prime}\right)=k\left(\underline{X}^{\prime}, \underline{X}\right)
$$

- for any $N \in \mathbb{N}$ and any $\left(\underline{X}_{1}, \ldots, \underline{X}_{N}\right) \in \mathcal{X}^{N}$,

$$
\boldsymbol{K}=\left[k\left(\underline{X}_{i}, \underline{X}_{j}\right)\right]_{1 \leq i, j \leq N}
$$

is positive semi-definite, i.e. $\forall u \in \mathbb{R}^{N}$

$$
u^{\top} \boldsymbol{K} u=\sum_{1 \leq i, j \leq N} u^{(i)} u^{(j)} k\left(\underline{X}_{i}, \underline{X}_{j}\right) \geq 0
$$

or equivalently all the eigenvalues of $\boldsymbol{K}$ are non-negative.

- The matrix $\boldsymbol{K}$ is called the Gram matrix associated to $\left(\underline{X}_{1}, \ldots, \underline{X}_{N}\right)$.


## 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

$$
k\left(\underline{X}, \underline{X}^{\prime}\right)=\left\langle\phi(\underline{X}), \phi\left(\underline{X}^{\prime}\right)\right\rangle_{\mathbb{H}}
$$

- the reproducing property holds, i.e. for any $h \in \mathbb{H}$ and any $\underline{X} \in \mathcal{X}$

$$
h(\underline{X})=\langle h, k(\underline{X}, \cdot)\rangle_{\mathbb{H}} .
$$

- 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.
- Rk: if $k\left(\underline{X}, \underline{X}^{\prime}\right)=\phi^{\prime}(\underline{X})^{\top} \phi^{\prime}\left(\underline{X}^{\prime}\right)$ with $\phi^{\prime}: \mathcal{X} \rightarrow \mathbb{R}^{p}$ then
- $\mathbb{H}$ can be chosen as $\left\{\underline{X} \mapsto \phi^{\prime}(\underline{X})^{\top} \beta, \beta \in \mathbb{R}^{p}\right\}$ and $\left\|\underline{X} \mapsto \phi^{\prime}(\underline{X})^{\top} \beta\right\|_{\mathbb{H}}^{2}=\|\beta\|_{2}^{2}$.
- $\phi\left(\underline{X}^{\prime}\right): \underline{X} \mapsto \phi^{\prime}(\underline{X})^{\top} \phi^{\prime}\left(\underline{X}^{\prime}\right)$.


## Separable Kernel

- For any function $\Psi: \mathcal{X} \rightarrow \mathbb{R}, k\left(\underline{X}, \underline{X}^{\prime}\right)=\Psi(\underline{X}) \Psi\left(\underline{X}^{\prime}\right)$ is PDS.


## Kernel Stability

- For any PDS kernels $k_{1}$ and $k_{2}, k_{1}+k_{2}$ and $k_{1} k_{2}$ 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 $\sum_{n} a_{n} z^{n}$ with $a_{n} \geq 0$ and a convergence radius larger than $r, \sum_{n} a_{n} k^{n}$ is a PDS kernel.
- For any PDS kernel $k$, the renormalized kernel $k^{\prime}\left(\underline{X}, \underline{X^{\prime}}\right)=\frac{k\left(\underline{X}, \underline{X^{\prime}}\right)}{\sqrt{k(\underline{X}, \underline{X}) k\left(\underline{X}^{\prime}, \underline{X}^{\prime}\right)}}$ is a PDS kernel.
- Cauchy-Schwartz for $k$ PDS: $k\left(\underline{X}, \underline{X}^{\prime}\right)^{2} \leq k(\underline{X}, \underline{X}) k\left(\underline{X}^{\prime}, \underline{X}^{\prime}\right)$


## PDS Kernels

- Vanilla kernel:

$$
k\left(\underline{X}, \underline{X}^{\prime}\right)=\underline{X}^{\top} \underline{X}^{\prime}
$$

- Polynomial kernel:

$$
k\left(\underline{X}, \underline{X}^{\prime}\right)=\left(1+\underline{X}^{\top} \underline{X}^{\prime}\right)^{k}
$$

- Gaussian RBF kernel:

$$
k\left(\underline{X}, \underline{X^{\prime}}\right)=\exp \left(-\gamma\left\|\underline{X}-\underline{X}^{\prime}\right\|^{2}\right)
$$

- Tanh kernel:

$$
k\left(\underline{X}, \underline{X}^{\prime}\right)=\tanh \left(a \underline{X}^{\top} \underline{X}^{\prime}+b\right)
$$

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


## Representer Theorem

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

$$
\underset{h \in \mathbb{H}}{\operatorname{argmin}} L\left(h\left(\underline{X}_{1}\right), \ldots, h\left(\underline{X}_{n}\right)\right)+\Phi(\|h\|)
$$

admits only solutions of the form

$$
\sum_{i=1}^{n} \alpha_{i}^{\prime} k\left(\underline{X}_{i}, \cdot\right)
$$

- Examples:
- (kernelized) SVM
- (kernelized) Penalized Logistic Regression (Ridge)
- (kernelized) Penalized Regression (Ridge)


## Kernelized SVM

## Primal

- Constrained Optimization:

$$
\min _{f \in \mathbb{H}, \beta^{(0)}, s}\|f\|_{\mathbb{H}}^{2}+C \sum_{i=1}^{n} s_{i} \quad \text { with } \quad\left\{\begin{array}{l}
\forall i, Y_{i}\left(f\left(\underline{X}_{i}\right)+\beta^{(0)}\right) \geq 1-s_{i} \\
\forall i, s_{i} \geq 0
\end{array}\right.
$$

- Hinge loss:

$$
\min _{f \in \mathbb{H}, \beta^{(0)}}\|f\|_{\mathbb{H}}^{2}+C \sum_{i=1}^{n} \max \left(0,1-Y_{i}\left(f\left(\underline{X}_{i}\right)+\beta^{(0)}\right)\right)
$$

- Representer:

$$
\begin{aligned}
& \min _{\alpha^{\prime}, \beta^{(0)}} \sum_{i, j} \alpha_{i}^{\prime} \alpha_{j}^{\prime} k\left(\underline{X}_{i}, \underline{X}_{j}\right) \\
& \quad+C \sum_{i=1}^{n} \max \left(0,1-Y_{i}\left(\sum_{j} \alpha_{j}^{\prime} k\left(\underline{X}_{j}, \underline{X}_{i}\right)+\beta^{(0)}\right)\right)
\end{aligned}
$$

## Dual

- Dual: $\max _{\alpha \geq 0, \mu \geq 0} Q(\alpha, \mu) \Leftrightarrow \max _{0 \leq \alpha \leq C} \sum_{i} \alpha_{i}-\frac{1}{2} \sum_{i, j} \alpha_{i} \alpha_{j} Y_{i} Y_{j} k\left(\underline{X}_{i}, \underline{X}_{j}\right)$

Support Vector Machine with polynomial kernel

Decision region


Decision boundary


Support Vector Machine with Gaussian kernel

Decision region


Decision boundary


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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)


## CART



## 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...


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
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## 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)

$$
C(R, \bar{R})=\sum_{\underline{x}_{i} \in R} \bar{\ell}\left(y_{i}, y(R)\right)+\sum_{\underline{x}_{i} \in \bar{R}} \bar{\ell}\left(y_{i}, y(\bar{R})\right)
$$

- CART: Gini index (Classification)

$$
C(R, \bar{R})=\sum_{\underline{x}_{i} \in R} p(R)(1-p(R))+\sum_{\underline{x}_{j} \in \bar{R}} p(\bar{R})(1-p(\bar{R}))
$$

- C4.5: entropy based criterion (Information Theory)

$$
C(R, \bar{R})=\sum_{\underline{x}_{i} \in R} H(R)+\sum_{\underline{x}_{i} \in \bar{R}} H(\bar{R})
$$

- 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...)

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 restricted set of points (quantiles or fixed grid)
- 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.

- 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

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

- Example: AIC / CV.


## Examples of criterion satisfying this assumptions

- AIC type criterion:

$$
\sum_{i=1}^{n} \bar{\ell}\left(y_{i}, f_{\mathcal{L}\left(\underline{x}_{i}\right)}\left(\underline{x}_{i}\right)\right)+\lambda|\mathcal{T}|=\sum_{\mathcal{L} \in \mathcal{T}}\left(\sum_{\underline{x}_{i} \in \mathcal{L}} \bar{\ell}\left(y_{i}, f_{\mathcal{L}}\left(\underline{x}_{i}\right)\right)+\lambda\right)
$$

- Simple cross-Validation (with $\left(\underline{x}_{i}^{\prime}, y_{i}^{\prime}\right)$ a different dataset):

$$
\sum_{i=1}^{n^{\prime}} \bar{\ell}\left(y_{i}^{\prime}, f_{\mathcal{L}}\left(\underline{x}_{i}^{\prime}\right)\right)=\sum_{\mathcal{L} \in \mathcal{T}}\left(\sum_{\underline{x}_{i}^{\prime} \in \mathcal{L}} \bar{\ell}\left(y_{i}^{\prime}, f_{\mathcal{L}}\left(\underline{x}_{i}^{\prime}\right)\right)\right)
$$

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

CART

Decision region


## Decision boundary



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)


Random Forest


## AdaBoost




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## Ensemble Methods



## Ensemble Methods

- Averaging: combine several models by averaging (bagging, random forests,... )
- Boosting: construct a sequence of (weak) classifiers (XGBoost, LightGBM, CatBoost)
- 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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5 A Probabilistic Point of View

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## Empirical Risk Minimizer (ERM)

- For any loss $\ell$ and function class $\mathcal{S}$,

$$
\widehat{f}=\underset{f \in \mathcal{S}}{\operatorname{argmin}} \frac{1}{n} \sum_{i=1}^{n} \ell\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)=\underset{f \in \mathcal{S}}{\operatorname{argmin}} \mathcal{R}_{n}(f)
$$

- Key property:

$$
\mathcal{R}_{n}(\widehat{f}) \leq \mathcal{R}_{n}(f), \forall f \in \mathcal{S}
$$

- 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:

$$
\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right)
$$

## Probably Almost Correct Analysis

- Theoretical guarantee that

$$
\mathbb{P}\left(\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \leq \epsilon_{\mathcal{S}}(\delta)\right) \geq 1-\delta
$$

for a suitable $\epsilon_{\mathcal{S}}(\delta) \geq 0$.

- Implies:
- $\mathbb{P}\left(\mathcal{R}(\widehat{f})-\mathcal{R}\left(f^{\star}\right) \leq \mathcal{R}\left(f_{\mathcal{S}}^{\star}\right)-\mathcal{R}\left(f^{\star}\right)+\epsilon_{\mathcal{S}}(\delta)\right) \geq 1-\delta$
- $\mathbb{E}\left[\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right)\right] \leq \int_{0}^{+\infty} \delta_{\mathcal{S}}(\epsilon) d \epsilon$
- The result should hold without any assumption on the law $\boldsymbol{P}$ !


## A General Decomposition

- By construction:

$$
\begin{aligned}
\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) & =\mathcal{R}(\widehat{f})-\mathcal{R}_{n}(\widehat{f})+\mathcal{R}_{n}(\widehat{f})-\mathcal{R}_{n}\left(f_{\mathcal{S}}^{\star}\right)+\mathcal{R}_{n}\left(f_{\mathcal{S}}^{\star}\right)-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \\
& \leq \mathcal{R}(\widehat{f})-\mathcal{R}_{n}(\widehat{f})+\mathcal{R}_{n}\left(f_{\mathcal{S}}^{\star}\right)-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \\
& \leq\left(\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right)\right)-\left(\mathcal{R}_{n}(\widehat{f})-\mathcal{R}_{n}\left(f_{\mathcal{S}}^{\star}\right)\right)
\end{aligned}
$$

## Four possible upperbounds

- $\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \leq \sup _{f \in \mathcal{S}}\left(\left(\mathcal{R}(f)-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right)\right)-\left(\mathcal{R}_{n}(f)-\mathcal{R}_{n}\left(f_{\mathcal{S}}^{\star}\right)\right)\right)$
- $\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \leq \sup _{f \in \mathcal{S}}\left(\mathcal{R}(f)-\mathcal{R}_{n}(f)\right)+\left(\mathcal{R}_{n}\left(f_{\mathcal{S}}^{\star}\right)-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right)\right)$
- $\mathcal{R}(\hat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \leq \sup _{f \in \mathcal{S}}\left(\mathcal{R}(f)-\mathcal{R}_{n}(f)\right)+\sup _{f \in \mathcal{S}}\left(\mathcal{R}_{n}(f)-\mathcal{R}(f)\right)$
- $\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \leq 2 \sup _{f \in \mathcal{S}}\left|\mathcal{R}(f)-\mathcal{R}_{n}(f)\right|$
- Supremum of centered random variables!
- Key: Concentration of each variable...
- By construction, for any $f^{\prime} \in \mathcal{S}$,

$$
\mathcal{R}\left(f^{\prime}\right)=\mathcal{R}_{n}\left(f^{\prime}\right)+\left(\mathcal{R}\left(f^{\prime}\right)-\mathcal{R}_{n}\left(f^{\prime}\right)\right)
$$

A uniform upper bound for the risk

- Simultaneously $\forall f^{\prime} \in \mathcal{S}$,

$$
\mathcal{R}\left(f^{\prime}\right) \leq \mathcal{R}_{n}\left(f^{\prime}\right)+\sup _{f \in \mathcal{S}}\left(\mathcal{R}(f)-\mathcal{R}_{n}(f)\right)
$$

- 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:

$$
\mathcal{R}_{n}(f)=\frac{1}{n} \sum_{i=1}^{n} \ell^{0 / 1}\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)
$$

## Properties

- $\ell^{0 / 1}\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)$ are i.i.d. random variables in $[0,1]$.


## Concentration

$$
\begin{gathered}
\mathbb{P}\left(\mathcal{R}(f)-\mathcal{R}_{n}(f) \leq \epsilon\right) \geq 1-e^{-2 n \epsilon^{2}} \\
\mathbb{P}\left(\mathcal{R}_{n}(f)-\mathcal{R}(f) \leq \epsilon\right) \geq 1-e^{-2 n \epsilon^{2}} \\
\mathbb{P}\left(\left|\mathcal{R}_{n}(f)-\mathcal{R}(f)\right| \leq \epsilon\right) \geq 1-2 e^{-2 n \epsilon^{2}}
\end{gathered}
$$

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


## Theorem

- Let $Z_{i}$ be a sequence of ind. centered r.v. supported in $\left[a_{i}, b_{i}\right]$ then

$$
\mathbb{P}\left(\sum_{i=1}^{n} Z_{i} \geq \epsilon\right) \leq e^{-\frac{2 \epsilon^{2}}{\sum_{i=1}^{n}\left(b_{i}-a_{i}\right)^{2}}}
$$

- Proof ingredients:
- Chernov bounds:

$$
\mathbb{P}\left(\sum_{i=1}^{n} z_{i} \geq \epsilon\right) \leq \frac{\mathbb{E}\left[e^{\lambda} \sum_{i=1}^{n} z_{i}\right]}{e^{\lambda \epsilon}} \quad \leq \frac{\prod_{i=1}^{n} \mathbb{E}\left[e^{\lambda z_{i}}\right]}{e^{\lambda \epsilon}}
$$

- Exponential moment bounds: $\mathbb{E}\left[e^{\lambda z_{i}}\right] \leq e^{\frac{\lambda^{2}\left(b_{i}-a_{i}\right)^{2}}{8}}$
- Optimization in $\lambda$
- Prop:

$$
\mathbb{E}\left[e^{\lambda \sum_{i=1}^{n} z_{i}}\right] \leq e^{\frac{\lambda^{2} \sum_{i=1}^{n}\left(b_{i}-a_{i}\right)^{2}}{8}}
$$

## Theorem

- Let $Z_{i}$ be a sequence of independent centered random variables supported in [ $a_{i}, b_{i}$ ] then

$$
\mathbb{P}\left(\sum_{i=1}^{n} Z_{i} \geq \epsilon\right) \leq e^{-\frac{2 \epsilon^{2}}{\left.\sum_{i=1}^{n} b_{i}-a_{i}\right)^{2}}}
$$

- $Z_{i}=\frac{1}{n}\left(\mathbb{E}\left[\ell^{0 / 1}(Y, f(\underline{X}))\right]-\ell^{0 / 1}\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)\right)$
- $\mathbb{E}\left[Z_{i}\right]=0$ and $Z_{i} \in\left[\frac{1}{n}\left(\mathbb{E}\left[\ell^{0 / 1}(Y, f(\underline{X}))\right]-1\right), \frac{1}{n} \mathbb{E}\left[\ell^{0 / 1}(Y, f(\underline{X}))\right]\right]$
- Concentration:

$$
\mathbb{P}\left(\mathcal{R}(f)-\mathcal{R}_{n}(f) \geq \epsilon\right) \leq e^{-2 n \epsilon^{2}}
$$

- By symmetry,

$$
\mathbb{P}\left(\mathcal{R}_{n}(f)-\mathcal{R}(f) \geq \epsilon\right) \leq e^{-2 n \epsilon^{2}}
$$

- Combining the two yields

$$
\mathbb{P}\left(\left|\mathcal{R}_{n}(f)-\mathcal{R}(f)\right| \geq \epsilon\right) \leq 2 e^{-2 n \epsilon^{2}}
$$

## Concentration

- If $\mathcal{S}$ is finite of cardinality $|\mathcal{S}|$,

$$
\begin{aligned}
& \mathbb{P}\left(\sup _{f}\left(\mathcal{R}(f)-\mathcal{R}_{n}(f)\right) \leq \sqrt{\frac{\log |\mathcal{S}|+\log (1 / \delta)}{2 n}}\right) \geq 1-\delta \\
& \mathbb{P}\left(\sup _{f}\left|\mathcal{R}_{n}(f)-\mathcal{R}(f)\right| \leq \sqrt{\frac{\log |\mathcal{S}|+\log (1 / \delta)}{2 n}}\right) \geq 1-2 \delta
\end{aligned}
$$

- 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 $\mathcal{S}$ is finite of cardinality $|\mathcal{S}|$, with proba greater than $1-2 \delta$

$$
\begin{aligned}
\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) & \leq \sqrt{\frac{\log |\mathcal{S}|+\log (1 / \delta)}{2 n}}+\sqrt{\frac{\log (1 / \delta)}{2 n}} \\
& \leq 2 \sqrt{\frac{\log |\mathcal{S}|+\log (1 / \delta)}{2 n}}
\end{aligned}
$$

- If $\mathcal{S}$ is finite of cardinality $|\mathcal{S}|$, with proba greater than $1-\delta$, simultaneously $\forall f^{\prime} \in \mathcal{S}$,

$$
\begin{aligned}
\mathcal{R}\left(f^{\prime}\right) & \leq \mathcal{R}_{n}\left(f^{\prime}\right)+\sqrt{\frac{\log |\mathcal{S}|+\log (1 / \delta)}{2 n}} \\
& \leq \mathcal{R}_{n}\left(f^{\prime}\right)+\sqrt{\frac{\log |\mathcal{S}|}{2 n}}+\sqrt{\frac{\log (1 / \delta)}{2 n}}
\end{aligned}
$$

Finite Class Case

## PAC Bounds

- If $\mathcal{S}$ is finite of cardinality $|\mathcal{S}|$, with proba greater than $1-2 \delta$

$$
\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \leq \sqrt{\frac{\log |\mathcal{S}|}{2 n}}+\sqrt{\frac{2 \log (1 / \delta)}{n}}
$$

- If $\mathcal{S}$ is finite of cardinality $|\mathcal{S}|$, with proba greater than $1-\delta$, simultaneously $\forall f^{\prime} \in \mathcal{S}$,

$$
\mathcal{R}\left(f^{\prime}\right) \leq \mathcal{R}_{n}\left(f^{\prime}\right)+\sqrt{\frac{\log |\mathcal{S}|}{2 n}}+\sqrt{\frac{\log (1 / \delta)}{2 n}}
$$

- Risk increases with the cardinality of $\mathcal{S}$.
- Similar issue in cross-validation!
- No direct extension for an infinite $\mathcal{S}$...


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Concentration of the Supremum of Empirical Losses

- Supremum of Empirical losses:

$$
\begin{aligned}
\Delta_{n}(\mathcal{S})\left(\underline{X}_{1}, \ldots, \underline{X}_{n}\right) & =\sup _{f \in \mathcal{S}} \mathcal{R}(f)-\mathcal{R}_{n}(f) \\
& =\sup _{f \in \mathcal{S}}\left(\mathbb{E}\left[\ell^{0 / 1}(Y, f(\underline{X}))\right]-\frac{1}{n} \sum_{i=1}^{n} \ell^{0 / 1}\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)\right)
\end{aligned}
$$

## Properties

- Bounded difference:

$$
\left|\Delta_{n}(\mathcal{S})\left(\underline{X}_{1}, \ldots, \underline{X}_{i}, \ldots \underline{X}_{n}\right)-\Delta_{n}(\mathcal{S})\left(\underline{X}_{1}, \ldots \underline{X}_{i}^{\prime}, \ldots, \underline{X}_{n}\right)\right| \leq 1 / n
$$

## Concentration

$$
\mathbb{P}\left(\Delta_{n}(\mathcal{S})-\mathbb{E}\left[\Delta_{n}(\mathcal{S})\right] \leq \epsilon\right) \geq 1-e^{-2 n \epsilon^{2}}
$$

- 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

$$
\begin{aligned}
& \forall\left(\underline{X}_{i}\right)_{i=1}^{n},\left(\underline{X}_{i}^{\prime}\right)_{i=1}^{n} \in \mathbb{R}, \\
& \quad\left|g\left(\underline{X}_{1}, \ldots, \underline{X}_{i}, \ldots, \underline{X}_{n}\right)-g\left(\underline{X}_{1}, \ldots, \underline{X}_{i}^{\prime}, \ldots, \underline{X}_{n}\right)\right| \leq c_{i}
\end{aligned}
$$

## Theorem

- If $g$ is a bounded difference function and $\underline{X}_{i}$ are independent random variables then

$$
\begin{aligned}
& \mathbb{P}\left(g\left(\underline{X}_{1}, \ldots, \underline{X}_{n}\right)-\mathbb{E}\left[g\left(\underline{X}_{1}, \ldots, \underline{X}_{n}\right)\right] \geq \epsilon\right) \leq e^{\frac{-2 \epsilon^{2}}{\sum_{i=1}^{n} c_{i}^{2}}} \\
& \mathbb{P}\left(\mathbb{E}\left[g\left(\underline{X}_{1}, \ldots, \underline{X}_{n}\right)\right]-g\left(\underline{X}_{1}, \ldots, \underline{X}_{n}\right) \geq \epsilon\right) \leq e^{\frac{-2 \epsilon^{2}}{\sum_{i=1}^{n} c_{i}^{2}}}
\end{aligned}
$$

- Proof ingredients:
- Chernov bounds
- Martingale decomposition...


## McDiarmid Inequality

## Theorem

- If $g$ is a bounded difference function and $\underline{X}_{i}$ are independent random variables then

$$
\mathbb{P}\left(g\left(\underline{X}_{1}, \ldots, \underline{X}_{n}\right)-\mathbb{E}\left[g\left(\underline{X}_{1}, \ldots, \underline{X}_{n}\right)\right] \geq \epsilon\right) \leq e^{\frac{-2 \epsilon^{2}}{\sum_{i=1}^{c_{i}^{2}}}}
$$

- Using $g=\Delta_{n}(\mathcal{S})$ for which $c_{i}=1 / n$ yields immediately

$$
\mathbb{P}\left(\Delta_{n}(\mathcal{S})-\mathbb{E}\left[\Delta_{n}(\mathcal{S})\right] \geq \epsilon\right) \leq e^{\frac{-2 \epsilon^{2}}{\sum_{i=1}^{n} c_{i}^{2}}}=e^{-2 n \epsilon^{2}}
$$

- We derive then

$$
\mathbb{P}\left(\Delta_{n}(\mathcal{S}) \geq \mathbb{E}\left[\Delta_{n}(\mathcal{S})\right]+\epsilon\right) \leq e^{\frac{-2 \epsilon^{2}}{\sum_{i=1}^{n} c_{i}^{2}}}=e^{-2 n \epsilon^{2}}
$$

- It remains to upperbound

$$
\mathbb{E}\left[\Delta_{n}\right]=\mathbb{E}\left[\sup _{f \in \mathcal{S}} \mathcal{R}(f)-\mathcal{R}_{n}(f)\right]
$$

## Theorem

- Let $\sigma_{i}$ be a sequence of i.i.d. random symmetric Bernoulli variables (Rademacher variables):

$$
\mathbb{E}\left[\sup _{f \in \mathcal{S}}\left(\mathcal{R}(f)-\mathcal{R}_{n}(f)\right)\right] \leq 2 \mathbb{E}\left[\sup _{f \in \mathcal{S}} \frac{1}{n} \sum_{i=1}^{n} \sigma_{i} \ell^{0 / 1}\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)\right]
$$

## Rademacher complexity

- Let $B \subset \mathbf{R}^{n}$, 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 $B_{n}(\mathcal{S})=\left\{\left(\ell^{0 / 1}\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)\right)_{i=1}^{n}, f \in \mathcal{S}\right\}$.
- 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

$$
R_{n}(B)=\mathbb{E}\left[\sup _{b \in B} \frac{1}{n} \sum_{i=1}^{n} \sigma_{i} b_{i}\right] \leq \sqrt{\frac{2 M^{2} \log |B|}{n}}
$$

- If $B=B_{n}(\mathcal{S})=\left\{\left(\ell^{0 / 1}\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)\right)_{i=1}^{n}, f \in \mathcal{S}\right\}$, we have $M=1$ and thus

$$
R_{n}(B) \leq \sqrt{\frac{2 \log \left|B_{n}(\mathcal{S})\right|}{n}}
$$

- We obtain immediately

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

Finite Set Rademacher Complexity Bound

## Theorem

- With probability greater than $1-2 \delta$,

$$
\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \leq \mathbb{E}\left[\sqrt{\frac{8 \log \left|B_{n}(\mathcal{S})\right|}{n}}\right]+\sqrt{\frac{2 \log (1 / \delta)}{n}}
$$

- With probability greater than $1-\delta$, simultaneously $\forall f^{\prime} \in \mathcal{S}$

$$
\mathcal{R}\left(f^{\prime}\right) \leq \mathcal{R}_{n}\left(f^{\prime}\right)+\mathbb{E}\left[\sqrt{\frac{8 \log \left|B_{n}(\mathcal{S})\right|}{n}}\right]+\sqrt{\frac{\log (1 / \delta)}{2 n}}
$$

- This is a direct consequence of the previous bound.

Finite Set Rademacher Complexity Bound

## Corollary

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

$$
\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \leq \sqrt{\frac{8 \log |\mathcal{S}|}{n}}+\sqrt{\frac{2 \log (1 / \delta)}{n}}
$$

- If $\mathcal{S}$ is finite then with probability greater than $1-\delta$, simultaneously $\forall f^{\prime} \in \mathcal{S}$

$$
\mathcal{R}\left(f^{\prime}\right) \leq \mathcal{R}_{n}\left(f^{\prime}\right)+\sqrt{\frac{8 \log |\mathcal{S}|}{n}}+\sqrt{\frac{\log (1 / \delta)}{2 n}}
$$

- It suffices to notice that

$$
\left|B_{n}(\mathcal{S})\right|=\left|\left\{\left(\ell^{0 / 1}\left(Y_{i}, f\left(\underline{X}_{i}\right)\right)\right)_{i=1}^{n}, f \in \mathcal{S}\right\}\right| \leq|\mathcal{S}|
$$

Finite Set Rademacher Complexity Bound

- Same result with Hoeffding but with better constants!

$$
\begin{aligned}
\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) & \leq \sqrt{\frac{\log |\mathcal{S}|}{2 n}}+\sqrt{\frac{2 \log (1 / \delta)}{n}} \\
\mathcal{R}\left(f^{\prime}\right) & \leq \mathcal{R}_{n}\left(f^{\prime}\right)+\sqrt{\frac{\log |\mathcal{S}|}{2 n}}+\sqrt{\frac{\log (1 / \delta)}{2 n}}
\end{aligned}
$$

- Difference due to the crude upperbound of

$$
\mathbb{E}\left[\sup _{f \in \mathcal{S}}\left(\mathcal{R}(f)-\mathcal{R}_{n}(f)\right)\right]
$$

- Why bother?: We do not have to assume that $\mathcal{S}$ is finite!

$$
\left|B_{n}(\mathcal{S})\right| \leq 2^{n}
$$

## Outline

A Probabilistic Point of View- Parametric Conditional Density Modeling
- Non Parametric Conditional Density Modeling
- Generative Modeling
(6) Optimization Point of View
- Penalization
- (Deep) Neural Networks
- SVM
- Tree Based Methods
(7) Empirical Risk Minimization
- Empirical Risk Minimization
- ERM and PAC Bayesian Analysis
- Hoeffding and Finite Class
- McDiarmid and Rademacher Complexity
- VC Dimension
- Structural Risk Minimization
(8) References


## Back to the Bound

## Theorem

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

- Key quantity: $\mathbb{E}\left[\sqrt{\frac{8 \log \left|B_{n}(\mathcal{S})\right|}{n}}\right]$
- Hard to control due to its structure!


## A first data dependent upperbound

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

- Depends on the unknown P!


## Shattering Coefficient

## Shattering Coefficient (or Growth Function)

- The shattering coefficient of the class $\mathcal{S}, s(\mathcal{S}, n)$, is defined as

$$
\left.\left.s(\mathcal{S}, n)=\sup _{\left(\left(\underline{X}_{1}, Y_{1}\right), \ldots,\left(\underline{X}_{n}, Y_{n}\right)\right) \in(\mathcal{X} \times\{-1,1\})^{n}} \operatorname{Y}_{i}, f\left(\ell_{i}^{0 / 1}\left(\underline{X}_{i}\right)\right)\right)_{i=1}^{n}, f \in \mathcal{S}\right\} \mid
$$

- By construction, $\left|B_{n}(\mathcal{S})\right| \leq s(\mathcal{S}, n) \leq \min \left(2^{n},|\mathcal{S}|\right)$.


## A data independent upperbound

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

## Shattering Coefficient

## Theorem

- With probability greater than $1-2 \delta$,

$$
\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \leq \sqrt{\frac{8 \log s(\mathcal{S}, n)}{n}}+\sqrt{\frac{2 \log (1 / \delta)}{n}}
$$

- With probability greater than $1-\delta$, simultaneously $\forall f^{\prime} \in \mathcal{S}$,

$$
\mathcal{R}\left(f^{\prime}\right) \leq \mathcal{R}_{n}\left(f^{\prime}\right)+\sqrt{\frac{8 \log s(\mathcal{S}, n)}{n}}+\sqrt{\frac{\log (1 / \delta)}{2 n}}
$$

- Depends only on the class $\mathcal{S}$ !


## VC Dimension

- The VC dimension $d_{V C}$ of $\mathcal{S}$ is defined as the largest integer $d$ such that

$$
s(\mathcal{S}, d)=2^{d}
$$

- The VC dimension can be infinite!


## VC Dimension and Dimension

- Prop: If $\operatorname{span}(\mathcal{S})$ corresponds to the sign of functions in a linear space of dimension $d$ then $d V C \leq d$.
- VC dimension similar to the usual dimension.


## VC Dimension and Sauer's Lemma

## Sauer's Lemma

- If the VC dimension $d_{V C}$ of $\mathcal{S}$ is finite

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

- Cor.: $\log s(\mathcal{S}, n) \leq d_{V C} \log \left(\frac{e n}{d_{V C}}\right)$ if $n>d_{V C}$.


## VC Dimension and PAC Bounds

## PAC Bounds

- If $\mathcal{S}$ is of VC dimension $d_{V C}$ then if $n>d_{V C}$
- With probability greater than $1-2 \delta$,

$$
\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \leq \sqrt{\frac{8 d_{V c} \log \left(\frac{e n}{d_{V C}}\right)}{n}}+\sqrt{\frac{2 \log (1 / \delta)}{n}}
$$

- With probability greater than $1-\delta$, simultaneously $\forall f^{\prime} \in \mathcal{S}$,

$$
\mathcal{R}\left(f^{\prime}\right) \leq \mathcal{R}_{n}\left(f^{\prime}\right)+\sqrt{\frac{8 d_{V C} \log \left(\frac{e n}{d_{V C}}\right)}{n}}+\sqrt{\frac{\log (1 / \delta)}{2 n}}
$$

- Rk: If $d_{V C}=+\infty$ no uniform PAC bounds exists!


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## Countable Collection and Non Uniform PAC Bounds

## PAC Bounds

- Let $\pi_{f}>0$ such that $\sum_{f \in \mathcal{S}} \pi_{f}=1$
- With proba greater than $1-2 \delta$,

$$
\mathcal{R}(\widehat{f})-\mathcal{R}\left(f_{\mathcal{S}}^{\star}\right) \leq \sqrt{\frac{\log \left(1 / \pi_{f}\right)}{2 n}}+\sqrt{\frac{2 \log (1 / \delta)}{n}}
$$

- With proba greater than $1-\delta$, simultaneously $\forall f^{\prime} \in \mathcal{S}$,

$$
\mathcal{R}\left(f^{\prime}\right) \leq \mathcal{R}_{n}\left(f^{\prime}\right)+\sqrt{\frac{\log \left(1 / \pi_{f}\right)}{2 n}}+\sqrt{\frac{\log (1 / \delta)}{2 n}}
$$

- Very similar proof than the uniform one!
- Much more interesting idea when combined with several models...


## Models, Non Uniform Risk Bounds and SRM

- Assume we have a countable collection of set $\left(\mathcal{S}_{m}\right)_{m \in \mathcal{M}}$ 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 $f \in \mathcal{S}_{m}$,

$$
\mathcal{R}(f) \leq \mathcal{R}_{n}(f)+\mathbb{E}\left[\sqrt{\frac{8 \log \left|B_{n}\left(\mathcal{S}_{m}\right)\right|}{n}}\right]+\sqrt{\frac{\log \left(1 / \pi_{m}\right)}{2 n}}+\sqrt{\frac{\log (1 / \delta)}{2 n}}
$$

## Structural Risk Minimization

- Choose $\hat{f}$ as the minimizer over $m \in \mathcal{M}$ and $f \in \mathcal{S}_{m}$ of

$$
\mathcal{R}_{n}(f)+\mathbb{E}\left[\sqrt{\frac{8 \log \left|B_{n}\left(\mathcal{S}_{m}\right)\right|}{n}}\right]+\sqrt{\frac{\log \left(1 / \pi_{m}\right)}{2 n}}
$$

- Mimics the minimization of the integrated risk!


## SRM and PAC Bound

## PAC Bound

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

$$
\begin{aligned}
\mathcal{R}(\widehat{f}) \leq & \inf _{m \in \mathcal{M}} \inf _{f \in \mathcal{S}_{m}}\left(\mathcal{R}(f)+\mathbb{E}\left[\sqrt{\frac{8 \log \left|B_{n}\left(\mathcal{S}_{m}\right)\right|}{n}}\right]+\sqrt{\frac{\log \left(1 / \pi_{m}\right)}{2 n}}\right) \\
& +\sqrt{\frac{2 \log (1 / \delta)}{n}}
\end{aligned}
$$

- The SRM minimizer balances the risk $\mathcal{R}(f)$ and the upper bound on the estimation error $\mathbb{E}\left[\sqrt{\frac{8 \log \left|B_{n}\left(\mathcal{S}_{m}\right)\right|}{n}}\right]+\sqrt{\frac{\log \left(1 / \pi_{m}\right)}{2 n}}$.
- $\mathbb{E}\left[\sqrt{\frac{8 \log \left|B_{n}\left(\mathcal{S}_{m}\right)\right|}{n}}\right]$ can be replaced by an upper bound (for instance a VC based one)...


## Outline

(1)

Introduction Machine Learning

- Motivation
(2)
- Method or Models
- Interpretability
- Metric Choice
(3) $A$

A Better Point of View

- The Example of Univariate Linear Regression
- Supervised Learning
(4)

Risk Estimation and Method Choice

- Cross Validation
- Cross Validation and Test
- Cross Validation and Weights
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- Parametric Conditional Density Modeling
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Per View

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