### Statistical Learning

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### Índex

- 1 Introduction and scope
- Introduction
- Statistical modelling
- Modelling data
- Detailed notation
- 6 Bibliography





## Preliminary note

The material in these slides is strongly based on [1]. When other materials are used, they are cited accordingly.

Mathematical notation follows as good as it can a good practices proposal from the Beijing Academy of Artificial Intelligence.





## What to expect?

#### In this session we will discuss:

- Modelling data
- Models with independent and identically distributed (iid) data
- The modelling dilemma
- Linear models
- Multivariate normal models
- •





## How is data analyzed and used?

Statistical learning interpret the model and quantify the uncertainity of the data.

Machine learning (or *data mining* making predictions using large scale data.

The goals of modelling data are:

- to predict data, based on existing one;
- to discover unusual or interesting patterns in data.

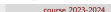




# Example of modelling data I

Imagine an unsupervised learning problem, with data represented by a vector  $\mathbf{x} = [x_1, \dots, x_p]^\mathsf{T}$ , a very general model is to assume that  $\mathbf{x}$  is the outcome of a random vector  $\mathbf{X} = [X_1, \dots, X_p]^\mathsf{T}$  with some unknown pdf f. The model can be refined by assuming a specific form of f.





# Example of modelling data II

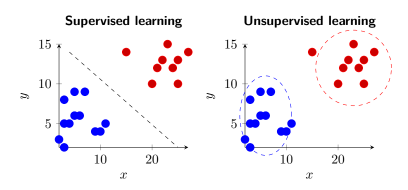


Figure 1: Supervised vs unsupervised ML





### Tools to model data

- Function approximation Model data with approximate and simple functions or maps.
- Optimization Given a set of feasible mathematical models to the data, we may need to find the optimal one by fitting or callibrating a function to observed data.
- Probability and Statistics Probability theory and statistical inference provides ways to quantify the uncertainity inherent in making predictions based on observed data.





### iid data

If we are given a sequence of data vectors  $x_1, \ldots, x_n$  one of the simplest possible models is to assume that the corresponding random vectors  $X_1, \ldots, X_n$  are independent and identically distributed (iid). We express this as:

$$X_1,\ldots,X_n \stackrel{iid}{\sim} f$$

meaning that the random vectors form an iid sample from a pdf f or sampling distribution Dist.

This is the same as saying that knowing about one variable does not provide information about another variable.





## Independent data models

In independent data models, the joint density of the random vectors  $X_1, \ldots, X_n$  is the *product* of the marginal ones:

$$f_{\mathbf{X}_1,\ldots,\mathbf{X}_n}(\mathbf{x}_1,\ldots,\mathbf{x}_n)=f(\mathbf{x}_1)\cdots f(\mathbf{x}_n)$$

The function g(x), the "model" for f(x) is usually specified up to a small number of parameters, corresponding to :

- $\mathcal{N}(\mu, \sigma^2)$
- Bin(n, p)
- Exp(λ)

The parameters are typically obtained from the data.





## Modeling dilemma

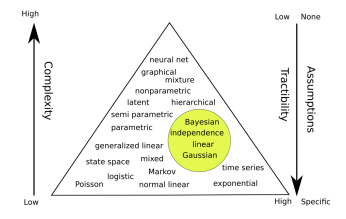


Figure 2: Complex models (very few of them) generally applicable but difficult to analyze. Simple models (a lot of options) very tractable but they do not describe well the data[1].

### Tradeoff

There exists a tradeoff between model tractability and applicability, as seen in Figure 2. Coming back to the example in page 6, the *training set*  $\tau = \{x_1, \dots, x_n\}$  is viewed as the outcome of n iid random variables  $X_1, \dots, X_n$  for some unknown pdf.

**Goal:** to learn or estimate *f* from the finite training set.





### Tradeoff vs risk

Imagine the **unsupervised learning** framewrok shown before. We can specify a class (a collection) of pdfs that we will call  $\mathcal{G}_p$ :

- We seek within  $\mathcal{G}$  the best approximation to the true model pdf  $f(\mathbf{x})$ , and we will call it  $g(\mathbf{x}|\Theta)$ .
- Such best approximation will minimize some calculated risk.

$$Loss(f(\mathbf{x}), g(\mathbf{x}|\Theta))) = \ln f(\mathbf{x}) - \ln g(\mathbf{x}|\Theta))$$

with expected value, this is, the risk as

$$\ell(g) = \mathbb{E} \ln \frac{f(\mathbf{X})}{g(\mathbf{X}|\Theta)} = \int f(\mathbf{x}) \ln \frac{f(\mathbf{x})}{g(\mathbf{x}|\Theta)} \, \mathrm{d}\mathbf{x}$$





### Train-Test-Validate

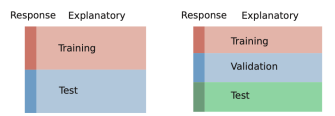


Figure 3: Sometimes we use the second set of data for model validation. Then we need to use a third one for testing.



### Train-Test-Validate

To compare the predictive performance of various learners in  $\mathcal{G}$ , as measured by the test loss,

- we can use the same fixed training set and test set for different learners, or
- if the overall data set is of modest size, we can perform the validation phase (model selection) on the training set only, using cross-validation.





# Polynomial regression. Original data.

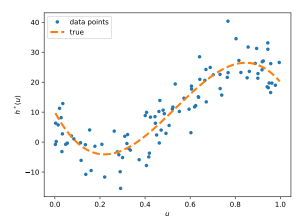


Figure 4: Training data and the optimal polynomial prediction function between the continuous and the optimal polynomial prediction function functi



# Polynomial regression. Fitting.

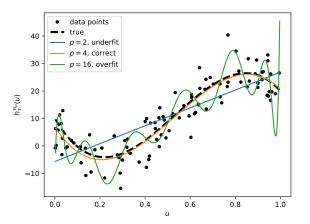


Figure 5: Fitted models for different orders of polynomial regressions.

# Polynomial regression. Error.

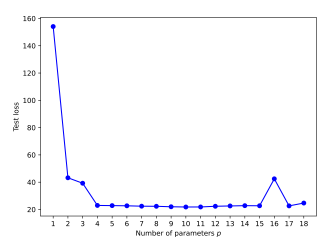


Figure 6: Fitted models for different orders of polynomial regressions.

## Polynomial regression. Cross validation.

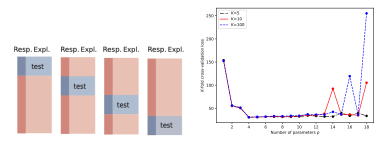


Figure 7: a) Example of four-fold cross-validation, representing four copies of the same data set. The data in each copy is partitioned into a training set (pink) and a test set (blue). Darker columns are the response variable; and lighter ones the explanatory variables. b) K-fold cross-validation for the polynomial regression.



# Some detailed notation (just for reference) I

Given an input or feature vector  $\mathbf{x}$ , ML aims at predicting an ouput or response variable vector  $\mathbf{y}$ . In particular, we search for a mathematical prediction function  $\mathbf{g}$  such that we can guess an approximation to  $\mathbf{y}$ ,  $\hat{\mathbf{y}}$ :

$$g: \mathcal{X} \to \mathcal{Y}$$
  
 $\mathbf{x} \mapsto \hat{\mathbf{y}} = g(\mathbf{x})$ 

#### Definition

Dataset  $S = \{z_i\}_{i=1}^n = \{(x_i, y_i)\}_{i=1}^n$  is sampled from a distribution  $\mathcal{D}$  over a domain  $\mathcal{Z} = \mathcal{X} \times \mathcal{Y}$ .

 $\mathcal X$  is the instance domain (a set),  $\mathcal Y$  is the label domain (a set), and  $\mathcal Z=\mathcal X\times\mathcal Y$  is the example domain (a set).



# Some detailed notation (just for reference) II

Usually,  $\mathcal{X}$  is a subset of  $\mathbb{R}^d$  and  $\mathcal{Y}$  is a subset of  $\mathbb{R}^{d_o}$ , where d is the input dimension,  $d_o$  is the output dimension.

n = #S is the number of samples. Without specification, S and n are for the training set.

- In regression problems, **y** is a vector of real values.
- In *classification* problems, y values lie within a finite set of c categories:  $y \in \{0, 1, \dots, c-1\}$ .

#### Definition

A hypothesis space is denoted by  $\mathcal{H}$ . A hypothesis function is denoted by  $f_{\theta}(\mathbf{x}) \in \mathcal{H}$  or  $f(\mathbf{x}; \theta) \in \mathcal{H}$  with  $f_{\theta} : \mathcal{X} \to \mathcal{Y}$ .

heta denotes the set of parameters of  $f_{ heta}$ .

If there exists a target function, it is denoted by  $f^*$  or  $f: \mathcal{X} \to \mathcal{Y}$  satisfying  $\mathbf{y}_i = f^*(\mathbf{x}_i)$  for  $i = 1, \dots, n$ .

# Some detailed notation (just for reference) III

A loss function, denoted by  $\ell:\mathcal{H}\times\mathcal{Z}\to\mathbb{R}_+:=[0,+\infty)$ , measures the difference (or error) between a predicted label and a true label, e.g.,  $L^2$  loss:

$$\ell(f_{\boldsymbol{\theta}}, \boldsymbol{z}) = \frac{1}{2}(f_{\boldsymbol{\theta}}(\boldsymbol{x}) - \boldsymbol{y})^2,$$

where  $\mathbf{z} = (\mathbf{x}, \mathbf{y})$ .  $\ell(f_{\theta}, \mathbf{z})$  can also be written as

$$\ell(f_{\boldsymbol{\theta}}(\boldsymbol{x}), \boldsymbol{y})$$

for convenience.

(In the case of a classification,  $\ell(f_{\boldsymbol{\theta}}, \boldsymbol{y}) = \mathbb{1}\{y \neq \hat{\boldsymbol{y}}\}$ )

We will see other useful loss functions ({em cross entropy} or *hinge* loss functions) later in this course.

It is unlikely that a mathematical function  $g \equiv f_{\theta} : \mathcal{X} \to \mathcal{Y}$  would be able to make accurate predictions of all possible pairs  $\mathcal{Z} = \mathcal{X} \times \mathcal{Y}$ .



# Some detailed notation (just for reference) IV

So, we use a probabilistic approach here to mpirical risk or training loss for a set  $S = \{(\mathbf{x}_i, \mathbf{y}_i)\}_{i=1}^n$  is denoted by  $L_S(\theta)$  or  $L_n(\theta)$  or  $R_n(\theta)$  or  $R_S(\theta)$ ,

$$L_{S}(\boldsymbol{\theta}) = \frac{1}{n} \sum_{i=1}^{n} \ell(f_{\boldsymbol{\theta}}(\mathbf{x}_{i}), \mathbf{y}_{i}). \tag{1}$$

The population risk or expected loss is denoted by  $L_{\mathcal{D}}(m{ heta})$  or  $R_{\mathcal{D}}(m{ heta})$ 

$$L_{\mathcal{D}}(\boldsymbol{\theta}) = \mathbb{E}_{\mathcal{D}}\ell(f_{\boldsymbol{\theta}}(\boldsymbol{x}), \boldsymbol{y}), \tag{2}$$

where  $\mathbf{z} = (\mathbf{x}, \mathbf{y})$  follows the distribution  $\mathcal{D}$ .

(In the case of a classification, we denote  $L_{\mathcal{D}}(g) \equiv L_{\mathcal{D}}(\theta) = \mathbb{P}_{\mathcal{D}}[f_{\theta}(\mathbf{x}) \neq \mathbf{y}]$  and we say that g is a classifier.)

Because we are interested in minimizing the risk in our prediction, we are looking for the best possible  $g^*$ : =  $\operatorname{argmin}_{\sigma} \mathbb{E}_{\mathcal{D}} \ell(f_{\theta}(\mathbf{x}), \mathbf{y})$ 



# Some detailed notation (just for reference) V

(In classification, we look for 
$$g^*(\mathbf{x}) = \underset{y \in \{0,1,\dots,c-1\}}{\operatorname{argmax}} \mathbb{P}[Y=y \mid X=x].)$$

#### **Theorem**

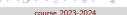
For the squared-error loss  $\ell(y, \hat{y}) = (y - \hat{y})^2$ , the optimal prediction function  $g^*$  is equal to the conditional expectation of Y given X = x.

which leads to write the random response Y as:

$$Y = g^*(\mathbf{x}) + \varepsilon(\mathbf{x})$$

Note that such random deviation satisfies  $\mathbb{E}\varepsilon(\mathbf{x})=0$ 







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Machine Learning & Pattern Recognition. Chapman & Hall/CRC, 2020.

