# 1. Introduction to Machine Learning

Jesper Armouti-Hansen

University of Cologne

December 10, 2018

#### Course content

- Introduction to Machine Learning
- 2 Linear methods for Classification
- Model Assessment and Selection
- 4 Linear methods for Regression
- 5 Moving beyond linearity
- 6 Tree-based methods
- Support Vector Machines

Format: mix of lectures, hands-on sessions, and case studies

#### Course website

jeshan49.github.io/eemp2/

### Today

- Lecture:
  - Motivation
- Tutorial:
  - Python basics
  - Introduction to NumPy
  - Getting started with Pandas
  - Plotting and visualization

# What is Machine Learning (ML)?

#### **General Definition**

[Machine Learning is the] field of study that gives computers the ability to **learn** without being explicitly programmed.

(Arthur Samuel, 1959)

#### More specific...

A computer program is said to learn from experience  ${\bf E}$  with respect to some task  ${\bf T}$  and some performance measure  ${\bf P}$ , if its performance on  ${\bf T}$ , as measured by  ${\bf P}$ , improves with experience  ${\bf E}$ .

■ (Tom Mitchell, 1997)

# Types of ML

#### **Supervised Learning**

- The task of learning a function that maps an input to an output based on example input-output pairs
- The learning method learns from a training sample consisting of a set of input-output obervations
- Depending on the nature of the output, the learning method should generalize from the training data to unseen data in order to either
  - give a good prediction of the output for new input if the output is quantitative
  - give a good classification of the output for new input if the output is qualitative
- Examples:
  - classifiy emails into "spam" or "ham" based on the content
  - predict the price of a house given a set of features

#### **Unsupervised Learning**

- We observe inputs but no outputs
- We can seek to understand the relationship between the variables or between the observations
- For example, in a market segmentation study we might observe multiple characteristics for potential customers and attempt to see if customers can be clustered into groups based on these characteristics
- If we have high-dimensional data, we might use unsupervised methods to project our inputs onto a lower dimensional space
- This can be a pre-processing step for a supervised learning method

### Reinforcement Learning

- The learning system, called an agent, can observe the environment, select and perform actions, and get a reward/punishment in return
- The agent then learns the best strategy in order to get the most reward over time
- The strategy defines what action the agent should choose when it is in a given situation
- Example: Deep-Q video

### Online vs. Batch Learning

- In batch learning, the learning method is incapable of learning incrementally
  - It must be trained using all available data
- Suppose we launch a method into production. Then, it is going to apply what it has already learned, but does not learn anymore: offline learning
- Thus, if we wish to update our model, we need to train a whole new version of it from scratch

- In online learning, you train the system incrementally by feeding it small data instances sequentially
- For example, if one wants a model that classifies spam/ham email, then it is probably a good option to have a model which can adapt to changes
- Online learning algorithms can also be used to train systems on huge data sets that cannot fit in one machine's memory
- An important parameter of the online learning algorithm is the learning rate
  - high: system will adapt quickly to new data, but quickly forget older
  - low: slow learning of new data, but will be less sensitive to noise in the new data

## Some Statistical Decision Theory

Let  $X \in \mathbb{R}^p$  denote a real valued random vector

i.e. the vector of inputs, features, predictors, or independent variables

Let  $Y \in \mathbb{R}$  be a real valued random output

• i.e. the response, target, or independent variable

Let Pr(X, Y) be their joint probability distribution

- We seek a function f(X) for predicting Y given values of the input X
- For this, we require a loss function L(Y, f(X)) that penalizes errors in prediction
- We will use the squared loss  $(Y f(X))^2$

Hence, we seek the function f(X) which minimizes

$$ESE(f) = E[(Y - f(X))^2]$$
(1)

$$= E[E[(Y - f(X))^{2}|X]]$$
 (2)

The solution is

$$f(x) = E[Y|X = x] \tag{3}$$

the conditional expectation, known as the regression function

- Thus, the best prediction of Y at any point X = x is the conditional mean
- Note:  $\varepsilon = Y f(X)$  is the irreducible error
  - even if we knew f(x), we would still make errors in prediction, since at each X = x there is typically a distribution of possible Y values.

Suppose we have estimated f(x) with  $\hat{f}(x)$  at the point x. Then we have

$$E[(Y - \hat{f}(X))^{2}|X = x] = E[(f(X) + \varepsilon - \hat{f}(X))^{2}|X = x]$$

$$= \underbrace{[f(X) - \hat{f}(X)]^{2}}_{\text{reducible}} + \underbrace{Var[\varepsilon]}_{\text{irreducible}}$$

We focus on techniques for estimating f with the aim of minimizing the reducible error.

The irreducible error provides an upper bound on the accuracy, but is almost always unknown.

### Estimating f - kNN regression

With our training data, we might attempt to directly apply the concept of E[Y|X=x] by asking, at each point x, for the average  $y_i$ .

Since we rarely have sufficient data points to do this, we can settle for:

$$\hat{f}(x) = Ave(y_i|x_i \in N_k(x)) \tag{4}$$

We perform two approximations in this approach:

- Expectation is approximated by averaging over sample data;
- Conditioning at a point is relaxed to conditioning on some region "close" to the target point

The k is called a *hyper* or *tuning parameter* - A parameter not learned from the learning procedure

## The Curse of Dimensionality

The kNN approach can work quite well for small p and large N

As p gets large, the nearest neighbors tend to be far away:

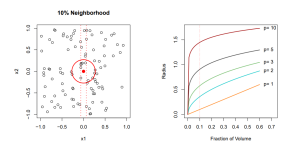


Figure: The radius of a sphere needed to capture a fraction of the volume of the data for different dimensions p (See ESL p. 23)

# Estimating f - Linear regression

How does linear regression fit into this framework?

• We simply assume that the regression function f(x) is approximately linear in its arguments

$$f(x) \approx x^{T} \beta \tag{5}$$

■ Plugging this linear model for f(x) into ESE in (1) and differentiating we can solve for  $\beta$ :

$$\beta = [E[XX^T]]^{-1}E[XY] \tag{6}$$

the least squares solution:

$$\hat{\beta} = (\mathbf{X}^T \mathbf{X})^{-1} \mathbf{X}^T \mathbf{y} \tag{7}$$

amounts to replacing the expectation by averages over the training data

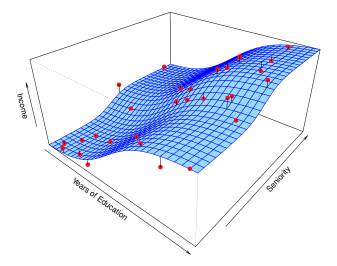


Figure: Simulated data: Income as a function of years of education and seniority. The blue surface represents the true underlying relationship. Red dots indicate observed values for 30 individuals (See ISLR p. 18)

## Parametric vs. Non-parametric Methods

Our goal: Apply a learning method in order to estimate f such that, for any observation (X, Y), we have  $Y \approx \hat{f}(X)$ .

Broadly speaking, we can decompose the learning methods into one of the following groups:

- Parametric methods
- 2 Non-parametric methods

The parametric methods share the following two step approach:

- $\blacksquare$  We make an assumption about the functional form of f
  - For example, if we assume f is linear, we only need to estimate p+1 coefficients as opposed to an arbitrary p-dimensional function
- Based on our chosen functional form, we choose a procedure to fit the model
  - For example, for the linear model, we (may) use the least squares procedure.

The method is parametric as it reduces the problem down to estimating a set of parameters.

Potential disadvantages?

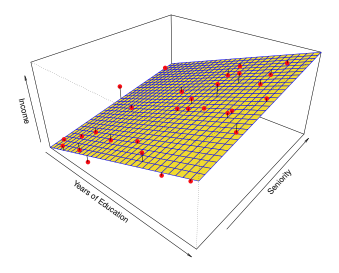


Figure: A linear model fit by least squares:  $income = \beta_0 + \beta_1 * education + \beta_2 * seniority$  (See ISLR p. 22)

#### Non-parametric methods:

- No explicit assumptions about the functional for of f
- Attempts to give an estimate of f close to observed data points subject to pre-specified constraints

#### Advantage:

lacktriangle Avoids making wrong functional form assumptions about f

#### Disadvantage:

Since the estimation problem is not reduced down to a set of parameters, a very large number of observations is required to obtain an accurate estimate.

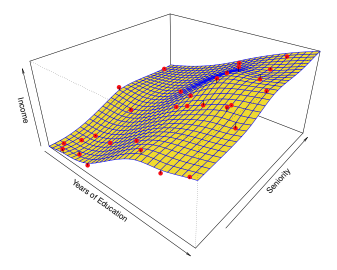


Figure: A smooth thin-plate spline fit to the same income data (See ISLR p. 23)

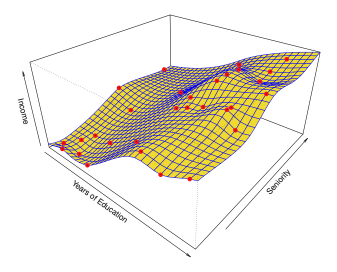


Figure: A rough thin-plate spline fit to the same income data (See ISLR p. 24)

# Prediction Accuracy vs. Model Interpretability

Why would we ever choose to use more restrictive models instead of a very flexible approach?

- As discussed, we might not have enough data
- If our goal includes model interpretability
  - Some of the models become so complex that understanding how any individual predictor is associated with the response becomes difficult
- We might overfit with highly flexible methods
- We often prefer a simple model involving fewer variables over a black-box model involving them all

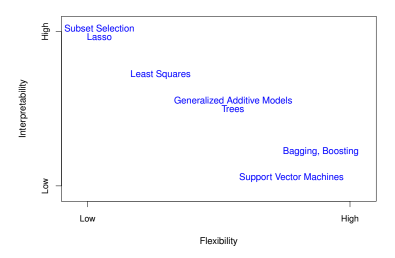


Figure: Tradeoff between flexibility and interpretability, using different learning methods (See ISLR p. 25)

### Assessing Model Accuracy

Why introduce many different learning approaches?

No Free Lunch: No one methods dominates all others over all possible data sets

Hence an important task is deciding on the best model for a given data set. To decide on a method, we need a metric to evaluate the quality of the fit

We could compute the average squared prediction error over Tr:

$$MSE_{Tr} = \frac{1}{N} \sum_{i=1}^{N} (y_i - \hat{f}(x_i))^2$$
 (8)

■ This may be biased toward more overfit models.

- What we would like to know is how our model, in general, will perform on new data
- If we are in a data-rich environment, we could have a designated hold-out or test set  $Te = \{x_i, y_i\}_1^M$  to estimate it:

$$MSE_{Te} = \frac{1}{M} \sum_{i=1}^{M} (y_i - \hat{f}(x_i))^2$$
 (9)

i.e., the test squared prediction error

- What if we don't have a large test set?
  - Perhaps we could use the training MSE!

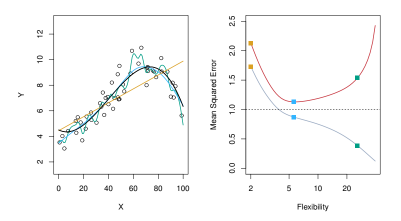


Figure: Simulated data: true f (black), linear regression line (orange), and two smoothing splines (blue and green) (See ISLR p. 31)

#### A more "linear" example:

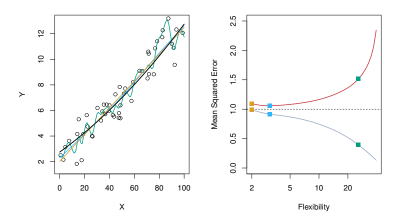


Figure: Simulated data: true f (black), linear regression line (orange), and two smoothing splines (blue and green) (See ISLR p. 33)

### A more "non-linear" example:

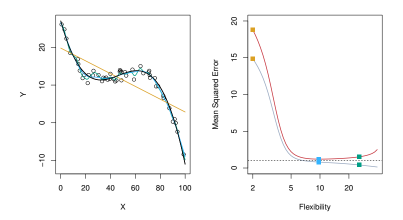


Figure: Simulated data: true f (black), linear regression line (orange), and two smoothing splines (blue and green) (See ISLR p. 33)

### The Bias-Variance Tradeoff

We saw that the test MSE tends to be U-shaped

- The shape is the result of two competing forces
- More formally, given  $x_0$ , the expected squared prediction error is given by

$$E[(y_0 - \hat{f}(x_0))^2] = E[y_0^2 + (\hat{f}(x_0))^2 - 2y_0 \hat{f}(x_0)]$$

$$= E[y_0^2] + E[(\hat{f}(x_0))^2] - E[2y_0 \hat{f}(x_0)]$$

$$= Var[y_0] + E[y_0]^2 + Var[\hat{f}(x_0)] + E[\hat{f}(x_0)]^2$$

$$- 2f(x_0)E[\hat{f}(x_0)]$$

$$= Var[\varepsilon] + \underbrace{Var[\hat{f}(x_0)]}_{\text{variance of } \hat{f}} + \underbrace{[f(x_0) - E[\hat{f}(x_0)]]^2}_{\text{bias}^2 \text{ of } \hat{f}}$$

- $E[(y_0 \hat{f}(x_0))^2]$  is the expected squared prediction error
  - i.e. the average test MSE if we repeatedly estimated f using a large number of training sets, and tested each at  $x_0$

A comparison of bias and variance in the three cases :

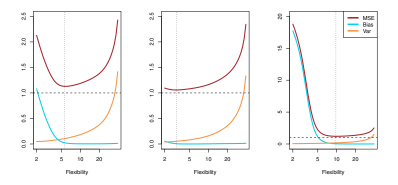


Figure: Squared bias (blue), variance (orange),  $Var(\varepsilon)$  (dashed), and test MSE (red) for the three data sets (See ISLR p. 36)

## The Classification Setting

Now Y is qualitative

lacktriangle e.g.  $\mathcal{C} = \{ \text{spam, ham} \}$  or  $\mathcal{C} = \{ 0, ..., 9 \}$ 

We wish to build a classifier C(X) that assigns a class label from  $\mathcal C$  to a future unlabeled observation X

- Suppoe C contains K elements numbered 1, ..., K
- Let  $p_k(x) = Pr(Y = k | X = x), k = 1, ..., K$
- Suppose we knew the conditional probability of Y given X
- Then, the Bayes optimal classifier at x given by

$$C(x) = j \text{ if } p_j(x) = \max\{p_1(x), ..., p_K(x)\}$$
 (10)

is optimal in the sense that it minimizes the expected one-zero loss:

$$E[\mathbb{I}(Y \neq C(X))] \tag{11}$$

## Estimating C - kNN classification

The Bayes classifier produces the lowest possible test error rate, called *Bayes error rate* 

$$1 - E[\max_{j} Pr(Y = j|X)] \tag{12}$$

We might attempt to apply kNN once again to

 $\blacksquare$  estimate the conditional distribution of Y given X

$$\hat{\rho}_j(x_0) = \frac{1}{K} \sum_{i \in \mathcal{N}_0} \mathbb{I}(y_i = j)$$
 (13)

classify a given observation to the class with highest estimated probability

$$\hat{C}(x) = j \text{ if } \hat{p}_j(x) = \max\{\hat{p}_1(x), ..., \hat{p}_K(x)\}$$
 (14)

Thus, kNN gives an estimate for the conditional probabilities as well as for the decision boundary

Note that the curse of dimensionality applies here too!

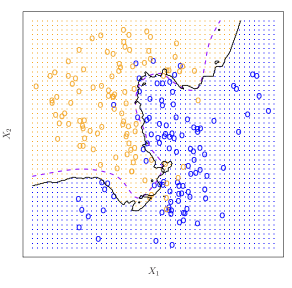


Figure: Bayes decision bounday (dashed), 10-NN decision boundary (black) (See ISLR p. 41)

#### A comparison of bias and variance in the three cases :

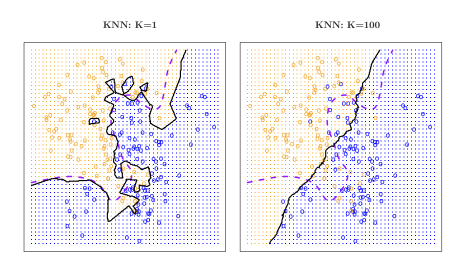


Figure: Bayes decision bounday (dashed), Left: 1-NN decision boundary (black), Right: 100-NN decision boundary (See ISLR p. 41)

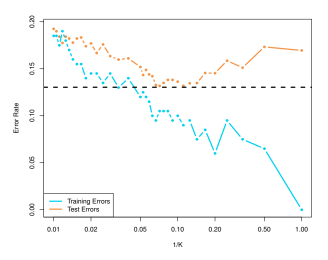


Figure: kNN training error rate (blue, 200 obs), test error rate (orange, 5,000 obs), Bayes error rate (dashed) (See ISLR p. 42)

- Some of the models we will consider build structured models for C(x)
  - e.g. support vector machines
- However, we will also build structured models for representing  $p_k(x)$ 
  - e.g. logistic regression and generalized additive models

# **Tutorial**

#### References

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