# Machine Learning for Education Policies

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# Intro

# General introduction to machine learning (ML)

ML is the sub-field of computer science that gives computers the ability to learn without being explicitly programmed, Arthur Samuel, 1959



# General introduction to machine learning (ML)

There are two cultures in the use of statistical modeling to reach conclusions from data. One assumes that the data are generated by a given stochastic data model. The other uses algorithmic models and treats the data mechanism as unknown. [...] If our goal as a field is to use data to solve problems, then we need to move away from exclusive dependence on data models and adopt a diverse set of tools, **Leo Breiman, 2001** 



# Key features of ML

- ML explores the study and construction of algorithms that can learn from and make predictions on data
- ML algorithms overcome following strictly static program instructions
- ML models do not make any assumptions about the data generating model (model free)
- The algorithms build a mathematical model from a set of data (data-driven models)

### What is learning?

A computer program is said to learn from experience E with respect to some class of tasks T and performance measure P if its performance at tasks in T, as measured by P, improves with experience E, **Tom M. Mitchell**, 1997

- E: experience is a set of inputs, colloquially a data set
- *T*: the class of tasks can be predictive, clustering, dimensionality reduction, anomaly detection, etc.
- P: performance measure that varies based on the task being tackled

E.g. Use data on students' SES, background and abilities to predict their financial literacy scores

# Inductive algorithms

### **Traditional Programming**



### **Machine Learning**



Slide credit: Pedro Domingos

### When is ML used?



- Humans expertise does not exist (navigating on Mars)
- Humans can't explain their expertise (speech recognition)
- Models are based on huge amount of data (genomics)
- Models must be customized (targeted policies)

# Supervised vs Unsupervised ML

Classification Regression Supervised Learning to predict. Unsupervised Learning to organize and represent. Dimensionality Clustering Reduction

### ML in social sciences

- Following Mullainathan and Speiss (2017 JEP) four main branches of applications:
  - 1. ML for causal inference (SL)
  - 2. ML for policy prediction (SL)
  - 3. ML to test theory (SL)
  - 4. ML for creation of new data sources (mostly UL)
- The focus will be on (1), (2)
- A brief overview on the some packages and functionalities for ML in R will be provided

Nutshell

Supervised Machine Learning in a

# Supervised Machine Learning (SML)

- The common denominator of SML algorithms is that they take an information set  $X_{N\times P}$  and map it to a vector of outputs y
- The functional form of this relationship is very **flexible** and gets updated by evaluating a loss function in two steps:
  - 1. pick the best loss-minimizing function  $f(\cdot)$ :

$$argmin \sum_{i=1}^{N} L(f(x_i), y_i)$$
 over  $f(\cdot) \in F$  s.t.  $R(f(\cdot)) \le c$ 

2. estimate the optimal level of complexity using empirical tuning through cross-validation

### Why is ML tailored for prediction?

• Take a generic loss function such as the MSE of prediction:

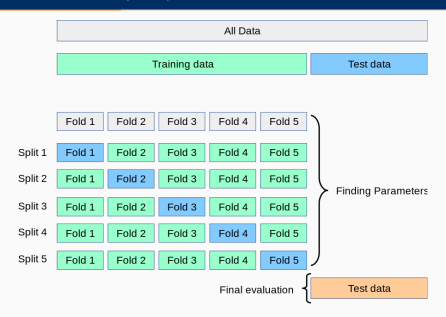
$$E_{\mathcal{D}}[(y - \hat{f}(x))^{2}] = \underbrace{E_{\mathcal{D}}[(E_{\mathcal{D}}[\hat{y_{0}}] - \hat{f}(x))^{2}]}_{\text{Variance}} + \underbrace{(E_{\mathcal{D}}[\hat{y_{0}}] - y)^{2}}_{\text{Bias}^{2}}$$

$$\underbrace{\text{Total Error}}_{\text{Variance}}$$
Variance

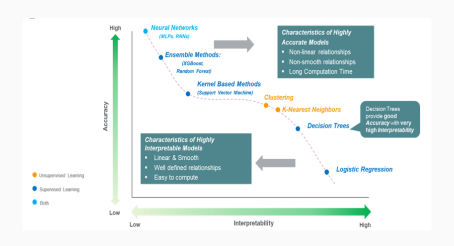
**Model Complexity** 

- By fixing the bias to be zero, the OLS regression rules out the possibility of this trade-off
- Impossibly to tune in a data-driven way the model with unbiased methods

### How is model complexity chosen?



# Interpretability vs Accuracy



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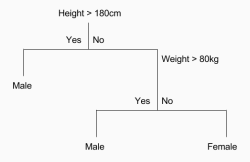
**Trees** 

Classification and Regression

### **Decision Trees**

### Definition 1 (Decision Tree)

Decision tree learning uses a decision tree (as a predictive model) to go from observations about an item (represented in the branches) to conclusions about the item's target value (represented in the leaves)



# Classification and Regression Trees

### Definition 2 (CART)

The CART methodology - introduced by Breiman, Friedman, Olshen and Stone in 1984 - is an algorithm for construction of binary trees, or trees where each node is splitted in only two branches

- Classification tree analysis is when the predicted outcome is the class to which the data belongs (e.g. *qualitative* outcomes)
- Regression tree analysis is when the predicted outcome can be considered a real number (e.g. quantitative outcomes)

CART is the basis for other algorithms that generate more complex trees. It is divided into two phases:

- 1. Generation of the tree
- 2. Pruning of the tree

# 1. Generation of a Regression tree

#### Generation of a tree:

- 1. Splitting of the predictor space (set of possible values for  $X_1, X_2, ..., X_p$ ) into J distinct and non-overlapping regions,  $R_1, R_2, ..., R_J$
- 2. Predict Y conditional on realization of  $X_j$  in each region  $R_j$  using the sample mean in that region

The construction of the regions  $R_1, R_2, ..., R_J$  (high-dimensional rectangles) proceeds by finding boxes  $R_1, R_2, ..., R_J$  that minimize the MSE given by:

$$\sum_{j=1}^{J} \sum_{i \in R_j} (y_i - \hat{y}_{R_j})^2$$

where  $-\hat{y}_{R_i}$  is the mean response for the training obs withing the j-th box

# Binary splitting

- Computationally infeasible to consider every possible partition of feature space
- 2. Top-down approach for the recursive binary splitting
  - a. Select a predictor  $X_i$  and a cut point s s.t.:

$$R_1(j,s) = \{X|X_j \le s\} \text{ and } R_2(j,s) = \{X|X_j > s\}$$

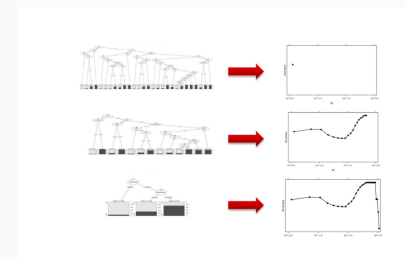
minimizes:

$$\sum_{i: x_i \in R_1(j,s)} (y_i - \hat{y}_{R_1})^2 + \sum_{i: x_i \in R_2(j,s)} (y_i - \hat{y}_{R_2})^2$$

- Repeat the process onto the two previously identified regions, so to minimize the MSE more
- Do it for all predictors and then choose the predictor and cut-point such that the resulting tree has the lowest MSE

# 2. Pruning of the tree (1)

Too complex trees lead to data overfitting



# 2. Pruning of the tree (2)

- · Two ways out:
  - 1. Split until the decrease in the MSE exceeds some threshold
  - 2. Grow a very large tree T and then prune it back to obtain a sub-tree
- This second strategy is implemented by minimizing:

$$\sum_{m=1}^{|\mathbb{T}|} \sum_{i:x_i \in R_m} (y_i - \hat{y}_{R_m})^2 + \alpha |\mathbb{T}|$$

where  $|\mathbb{T}|$  indicates the number of nodes of the tree  $\mathbb{T}$ ,  $R_m$  is the rectangle corresponding to the m-th terminal node and  $\alpha$  is a non-negative tuning parameter chosen by Cross-Validation

### **Classification Trees**

- Classification Trees are similar to Regression Trees except they are used to predict a qualitative response
- The main difference is that instead of minimizing the MSE it is used the Classification Error Rate

$$MSE \rightarrow CER$$

 CER is the fraction of training obs. in a region that do not belong to the most common class

$$CER = 1 - \max_{k} (\hat{p}_{m,k})$$

where  $\hat{p}_{m,k}$  represents the proportion of training obs. in the m-th region that are from the k-th class

# Impurity measure: Entropy and Information Gain

- Entropy and Information Gain
  - Definition: degree of disorder of our dataset  $\Omega$ : if we define by  $F_1$  and  $F_2$  the fraction of observations  $\Omega$  classified with "1" and "2", the entropy of the entire system S is defined as the following function H(S):

$$H(S) = -F_1 log F_1 - F_2 log F_2$$

• Respect to the J subclasses entropy is defined as:

$$H(S) = -\sum_{j=1}^{J} F_{j} log F_{j}$$

 The concept of information gain is a formalization of the entropic gain obtained through a partition of the data:

$$G = H(S) - H(S,s)$$
 where  $H(S,s) = H'(S)$ 

### Pros and Cons of CART

· Strengths and weaknesses of the CART methodology

#### · Pros:

- CART results are invariant under monotone transformations of the independent variables;
- CART can use the data set with a complex structure have been developed to be able to detect the dominant structures of the data;
- 3. CART are extremely robust to outliers;
- CART can use linear combinations of variables to make the split: no need to discretize continuous variables

#### Cons:

- 1. We don't use all the data (cross-validation);
- Every time our algorithm chooses a split, it chooses the best split in that exact moment (no bigger picture) → greedy algorithm

# Application of the CART methodology in R

Machine Learning and Causality Using CART to estimate heterogeneous causal effect

# **Machine Learning and Causality**

### · Econometrics/ Statistics/ Social Science

- Formal theory of causality
  - Potential outcomes methods (Rubin) maps onto economic approaches
- Well-developed and widely used tools for estimation and inference of causal effect in experimental and observational studies
  - Used by social science, policy-makers, development organizations, medicine, business, experimentation
- Weaknesses
  - · Non-parametric approaches fail with many covariates
  - · Model selection unprincipled

### Motivations

- · Experiments and Data-Mining
  - Concerns about ex-post "data-mining"
    - In medicine, scholars are required to pre-specify analysis plan (similar in economic field experiments)
- How is it possible to deal with sets of treatment effects among subsets of the entire population?
- Estimate of treatment effect heterogeneity needed for optimal decision-making

### Definition 3 (Athey and Imbens, 2015; 2016)

- Estimating heterogeneity by features in causal effects in experimental or observational studies
- 2. Conduct inference about the magnitude of the differences in the treatment effects across subsets of the population

### Causal Inference Framework

- Causal inference in observational studies:
  - As we saw previously, assuming unconfoundedness to hold, we can treat observations as having come from a randomized experiment
  - Therefore we can define the conditional average treatment effect (CATE) as follows:

$$\tau(x) = E[Y_i(1) - Y_i(0)|X_i = x]$$

· The population average treatment effect then is:

$$\tau^p = E[Y_i(1) - Y_i(0)] = E[\tau(X_i)]$$

# Why is CATE important?

- There are a variety of reasons that researchers wish to conduct estimation and inference on  $\tau(x)$ :
  - 1. It my be used to assign future units to their optimal treatment (in presence of different levels of the treatment):

$$W_i^{opt} = max \, \tau(X_i)$$

2. If we don't pre-specify the sub-populations it can be the case that the overall effect is negative, but it can be positive on subpopulations, then:

$$W_i^{PTE} = \mathbf{1}_{\tau(X_i) > 0}$$

e.g.: treatment is a drug  $\rightarrow$  prescribe it just to those who benefit from it

# Using Trees to Estimate Causal Effects

### Athey and Imbens (2015; 2016) propose 3 different approaches:

- Approach I: Analyze two groups separately:
  - Estimate  $\hat{\mu}(1, x)$  using dataset where  $W_i$ =1
  - Estimate  $\hat{\mu}(0,x)$  using dataset where  $W_i$ =0
  - Preform within group cross-validation to choose tuning parameters
  - Predict  $\hat{\tau} = \hat{\mu}(1, x) \hat{\mu}(0, x)$

- Approach II: Estimate  $\mu(w, x)$  using just one tree:
  - Estimate  $\hat{\mu}(1,x)$  and  $\hat{\mu}(0,x)$  using just one tree
  - Preform within tree cross-validation to choose tuning parameters
    - Predict  $\hat{\tau} = \hat{\mu}(1, x) \hat{\mu}(0, x)$
    - Estimate is zero for x where tree does not split on w

### The CATE Transformation of the Outcome

- The authors' goal is to develop an algorithm that generally leads to an accurate approximation of  $\hat{\tau}$  the Conditional Average Treatment Effect.
  - 1. Ideally we would measure the quality of the approximation in terms of goodness of fit using the MSE:

$$Q^{infeas} = \frac{1}{N} \sum_{i=1}^{N} (Y_i(1) - Y_i(0) - \hat{\tau}(X_i))^2$$

2. We can address this problem of infeasibility by transforming the outcome using the treatment indicator  $W_i$  and e(X):

$$Y_i^* = Y_i^{\text{obs}} \cdot \frac{W_i - e(X_i)}{(1 - e(X_i)) \cdot e(X_i)}$$

3. Then:

$$E[Y_i^*|X_I=x]=\tau(x)$$

### How to estimate the In-Sample Goodness of fit?

• The ideal goodness of fit measure would be:

$$Q^{infeas}(\hat{\tau}) = \mathbb{E}[(\tau_i - \hat{\tau}(X_i))^2].$$

 A useful proxy that can be used for the goodness of fit measure is:

$$\mathbb{E}[\tau_i^2|X_i\in S_j]=\frac{1}{N}\sum_i\hat{\tau}(x_i)^2.$$

This leads to our In-sample goodness of fit function:

$$Q^{is} = -\frac{1}{N} \sum_{i} \hat{\tau}(x_i)^2.$$

### Transformed Outcome Tree Model

- · Approach 3:
  - 1. Model and Estimation
    - · Model Type: Tree structure
    - Estimator  $\hat{\tau}_i^{TOT}$ : sample average treatment effect within leaf
  - 2. Criterion function (for fixed tuning parameter  $\lambda$ )
    - · In-sample Goodness-of-fit function:

$$Q^{is} = -MSE = -\frac{1}{N} \sum_{i=1}^{N} (\hat{\tau}_i^{TOT})^2$$

· Structure and use of criterion:

$$Q^{crit} = Q^{is} - \lambda \times leaves$$

- Select member of set of candidate estimators that maximizes  $Q^{crit}$ , given  $\lambda$
- 3. Cross-validation approach
  - · Out-of-Sample Goodness-of-fit function:

$$Q^{\text{oos}} = -MSE = -\frac{1}{N} \sum_{i=1}^{N} (\hat{\tau}_i^{TOT} - Y_i^*)^2$$

• Approach: select tuning parameter  $\lambda$  with highest  $Q^{os}$ 

## Critique to the TOT approach

Transformation of the Outcome in a randomized set-up:

$$Y_i^* = Y_i^{\text{obs}} \cdot \frac{W_i - p}{(1 - p) \cdot p} = \begin{cases} \frac{1}{p} \cdot Y_i^{\text{obs}} & \text{if } W_i = 1\\ -\frac{1}{1 - p} \cdot Y_i^{\text{obs}} & \text{if } W_i = 0 \end{cases}$$

- Within a leaf the sample average of Y<sub>i</sub>\* is not the most efficient estimator of treatment effect
- The proportion of treated units within the leaf is not the same as the overall sample proportion
- We use a weighted estimator similar to the Hirano, Imbens and Ridder (2003) estimator

## Causal Tree Approach

• In details the Treatment Effect in a generic leaf  $X_j$  is:

$$\tau^{CT}(X_i) = \frac{\sum_{j:X_j \in \mathbb{X}_j} y_i^{obs} \cdot \frac{W_i}{\hat{e}(X_i)}}{\sum_{j:X_j \in \mathbb{X}_j} \frac{W_i}{\hat{e}(X_i)}} - \frac{\sum_{j:X_j \in \mathbb{X}_j} y_i^{obs} \cdot \frac{(1 - W_i)}{(1 - \hat{e}(X_i))}}{\sum_{j:X_j \in \mathbb{X}_j} \frac{(1 - W_i)}{(1 - \hat{e}(X_i))}}$$

· This estimator is a consistent estimator of:

$$au_{\mathbb{X}_i} = \mathbb{E}[Y_i(1) - Y_i(0)|X_i \in \mathbb{X}_j]$$

· The variance can be estimated the Neyman estimator:

$$\hat{\mathbb{V}}_{Neyman} = \frac{s_t^2}{N_t} + \frac{s_c^2}{N_c}$$

These two quantities can be estimated as:

$$s_{t,j}^{te,2} = \frac{1}{N_t - 1} \sum_{i:W_i = 1} (Y_i(1) - \overline{Y}_t^{obs})^2 = \frac{1}{N_t - 1} \sum_{i:W_i = 1} (Y_i - \overline{Y}_t^{obs})^2$$
  
$$s_{c,j}^{te,2} = \frac{1}{N_c - 1} \sum_{i:W_i = 0} (Y_i(0) - \overline{Y}_c^{obs})^2 = \frac{1}{N_c - 1} \sum_{i:W_i = 0} (Y_i - \overline{Y}_c^{obs})^2$$

### Attractive features of Causal trees

- 1. Can easily separate tree construction from treatment effect estimation
- 2. Tree constructed on training sample is independent of sampling variation in the test sample
- Holding tree from training sample fixed, can use standard methods to conduct inference within each leaf of the tree on test sample
- 4. Can use any valid method for treatment effect estimation, not just the methods used in training
- 5. Simulations run by the authors show that the Causal Tree Algorithm over-performs the ST, TT and TOT approaches

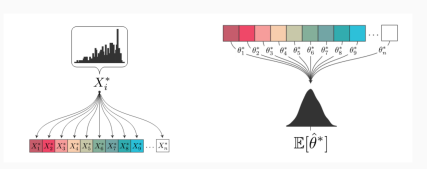
# Application of the CT methodology in R

# Extra on ML for heterogeneous effects

# Random forests

#### **Random Forest**

- RF: A Random Forest (Breiman, 2001) is a collection of fully grown CART. A
  Random Forest is a substantial transformation of the bagging method by
  introducing a collection of trees uncorrelated with each other.
  - 1. Bagging;
  - 2. Independence.



## 1. Bagging

- The techniques called bagging take shape bootstrap by the method. The term itself comes from bootstrap aggregation.
  - 1. Bradley Efron (1979)
  - 2. Sample **X** of dimension *n*
  - 3. Estimate the parameter  $\theta$  by simulating **B** samples of the same abundance, obtained sampling by assuming **X** as if for the overpopulation of reference
- $\hat{f}(X) \mapsto B \text{ samples } X_1^*, ..., X_B^* \mapsto t(X_1^*), ..., t(X_B^*)$
- · Bagging estimator

$$\hat{t}_{bag}(X) = \frac{1}{B} \sum_{b=1}^{B} t(X_b^*)$$

### 2. Independence

· If we develop the *bagged* variance estimator  $\hat{t}^{B}_{bag}(X)$ :

$$Var(\hat{t}_{bag}^{B}(X)) = ar\left(\frac{1}{B}\sum_{i=1}^{B}t(X_{i})\right)$$
$$= \sigma^{2} \cdot \rho + \sigma^{2}\frac{1-\rho}{B}$$

.

- The idea behind the Random Forest is that we can significantly increase the benefits of bagging through a reduction in the correlation of trees
- Random selection mechanism to select m variables between the p total splitting variables

#### Random Forest Algorithm for Regression and Classification

- 1. For *b* that goes from 1 to B:
  - Draw a sample Z\* of N units through the bootstrap method from our starting datasets Ω;
  - Grow a tree of the random forest  $T_b$  repeating, recursively, the following steps for each terminal node of the tree until you reach the minimum number of nodes  $n_{min}$ 
    - 1.i Select *m* randomly variables between the *p* available variables;
    - Choose the best combination of variables used for the split between the m variables;
    - 1.iii Splitting the node into 2 children nodes.
- 2. Through the output of all the trees  $\{T_b\}^B$  we can proceed as follows:
  - Regression:  $\hat{t}_{rf}^B(X) = \frac{1}{B} \sum_{b=1}^B T_b(x)$ ;
  - Classification: we can think about prediction of the k-th class and the b-th tree of the random forest as  $\hat{C}_{rf}^{B,K}(x)$ . Where:  $\hat{C}_{ff}^{B,K}(x)$ =majority vote  $\{\hat{C}_b(x)^{B,K}\}$

### Pros and Cons of RF

#### · Pros:

- 1. There is no need to rework or transform the data before building the model. Data must not be normalized and this approach is particularly robust to outliers;
- If we have a lot of input variables, we must not do any variable selection in a prior stage to construction of the model because it will be the same Random Forest to identify what are the most useful variables.;
- 3. Many trees are built through random mechanisms and therefore every tree is actually an independent model that does not bring the model to an overfitting.

#### · Cons:

- 1. Strong data dependency
- 2. Lower interpretability
- 3. Higher computational costs

# Application of the RF approach in R

## Extra on RF