## Machine Learning: A Very Brief Overview

A. Colin Cameron U.C.-Davis

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#### Introduction

- Machine learning is used to make predictions.
- The term machine learning is used because the machine learns from past data, rather than using models specified by knowledgable experts.
- For example, initial efforts for computer-based language translation used rules of grammar. Now machine learning predictions are used instead.
- There are many different machine learning algorithms.
- In many settings they perform much better than previous prediction methods.

### Regression

- Suppose we want to predict y based on potential variables  $x_1, ...., x_p$ .
- Traditional economics methods would use some combination of
  - specify a model such as a linear regression model and include only those variables that previous studies suggest are releavnt
  - possibly then drop variables that are statistically significant.
- Machine learning methods instead
  - specify a vary flexible nonlinear model
  - use methods that reduce the variability of the prediction by allowing some bias
  - use out-of-sample prediction to choose the best model and guard against in-sample overfitting.

#### Overview

- Terminology
- Model selection especially cross-validation.
- Variance-bias trade-off and shrinkage (LASSO and Ridge)
- Neural nets
- Regression trees and random forests
- Other Methods
- Classification
- Unsupervised learning (cluster analysis)
- Prediction for economics
- Causal Inference
- References



### 1. Terminology

- The term **machine learning** is used because the machine (computer) figures out from data the model  $\hat{y} = \hat{f}(\mathbf{x})$ 
  - compared to a modeler who e.g. specifies  $\mathbf{x}$  and  $y = \mathbf{x}'\boldsymbol{\beta} + u$ .
- The data may be big or small
  - ightharpoonup typically dim(m x) is large but n can be small or large.

# Terminology (continued)

#### Supervised learning = Regression

- We have both outcome y and regressors (or features) x
- ▶ 1. **Regression**: *y* is continuous
- 2. Classification: y is categorical.

#### Unsupervised learning

- We have no outcome y only several x
- ➤ 3. Cluster Analysis: e.g. determine five types of individuals given many psychometric measures.
- Focus on 1. as this is most used by economists.
- A lot of machine learning is actually used for 2.
  - ▶ license plate recognition, Google translate, ....

# Terminology (continued)

- Consider two types of data sets
  - ▶ 1. training data set (or estimation sample)
    - used to fit a model.
  - 2. test data set (or hold-out sample or validation set)
    - ★ additional data used to determine how good is the model fit
    - $\star$  a test observation  $(\mathbf{x}_0, y_0)$  is a previously unseen observation.

#### 2. Model selection

- Machine learners choose x's by using predictive ability
  - ▶ often mean squared error MSE =  $\frac{1}{n} \sum_{i=1}^{n} (y_i \widehat{y}_i)^2$ .
- Problem: models "overfit" within sample.
- Solution 1:
  - use an in-estimation-sample prediction with penalty for overfitting
    - $\star$  e.g.  $\bar{R}^2$ , AIC, BIC, Mallows Cp
- Solution 2:
  - use out-of-estimation sample prediction (cross-validation)
    - \* new to econometrics
    - ★ can apply to other loss functions and not just MSE.



#### K-fold cross-validation is standard method

- ullet K-fold cross-validation (standard choices are K=5 and K=10)
  - ▶ split data into K mutually exclusive folds of roughly equal size
  - for j = 1, ..., K fit using all folds but fold j and predict on fold j
- The following shows case K=5

	Fit on folds	Test on fold
j = 1	2,3,4,5	$1  o MSE_{(1)}$
j=2	1,3,4,5	$2 \rightarrow MSE_{(2)}$
<i>j</i> = 3	1,2,4,5	$3 \rightarrow MSE_{(3)}$
j = 4	1,2,3,5	$4 \rightarrow MSE_{(4)}$
<i>j</i> = 5	1,2,3,4	$5 \rightarrow MSE_{(5)}$

• The K-fold CV estimate is

$$CV_K = \frac{1}{K} \sum_{i=1}^K MSE_{(j)}$$
, where  $MSE_{(j)}$  is the MSE for fold  $j$ .

• Choose the model with smallest  $CV_K$ .



### 3. Bias-Variance Trade-off and Shrinkage Estimation

- The goal is minimize MSE = Variance + Bias-squared.
- More flexible models have
  - less bias (good) and more variance (bad).
  - this trade-off is fundamental to machine learning.
- Shrinkage reduces variance and may offset increased bias.
  - e.g.  $\widehat{\beta} = 0$  has reduced variance to zero.

## Shrinkage Methods: Ridge Regression

- Shrinkage estimators minimize RSS (residual sum of squares)
  - but with a penalty for model size
  - this shrinks parameter estimates towards zero.
- ullet The ridge estimator  $\widehat{oldsymbol{eta}}_{\lambda}$  of  $oldsymbol{eta}$  minimizes

$$Q_{\lambda}(\beta) = \sum_{i=1}^{n} \{ y_i - (\beta_1 x_{1i} + \dots + \beta_p x_{pi}) \}^2 + \lambda \sum_{j=1}^{p} \beta_j^2$$

- where  $\lambda \geq 0$  is a tuning parameter to be determined
- Equivalently minimize  $\sum_{i=1}^{n} \{ y_i (\beta_1 x_{1i} + \dots + \beta_p x_{pi}) \}^2$  subject to  $\sum_{i=1}^{p} \beta_i^2 \leq s$ .
- Typically first standardize x's to have mean zero and variance 1.

### Shrinkage Methods: LASSO

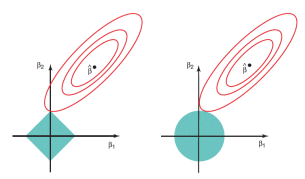
- Instead of squared penalty use absolute penalty.
- ullet The Least Absolute Shrinkage and Selection (LASSO) estimator  $\widehat{oldsymbol{eta}}_{\lambda}$  of  $oldsymbol{eta}$  minimizes

$$Q_{\lambda}(\beta) = \sum_{i=1}^{n} \{ y_i - (\beta_1 x_{1i} + \dots + \beta_p x_{pi}) \}^2 + \lambda \sum_{j=1}^{p} |\beta_j|$$

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- The acronym LASSO is because
  - it sets some  $\beta's$  to zero and shrinks others towards zero.

# LASSO versus Ridge (key figure from ISL)

- Two regressors: ellipses are the residual sum of squares for different values of  $\beta_1$  and  $\beta_2$  and the green squares are constraints
- Estimate is ellipse that just satisfies the constraint
  - ▶ LASSO is likely to be at a corner where some coefficients are zero.



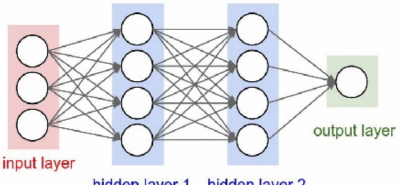
**FIGURE** 6.7. Contours of the error and constraint functions for the lasso (left) and ridge regression (right). The solid blue areas are the constraint regions,  $|\beta_1| + |\beta_2| \le s$  and  $\beta_1^2 + \beta_2^2 \le s$ , while the red ellipses are the contours of the RSS.

# 4. Neural Networks (deep learning)

- A neural network involves a series of nested regressions.
- ullet A single hidden layer neural network explaining y by  ${f x}$  has
  - ▶ y depends on z's (a hidden layer)
  - **z**'s depend on **x**'s.
- ullet A neural network with two hidden layers explaining y by  ${\bf x}$  has
  - ightharpoonup y depends on  $\mathbf{w}'s$  (a hidden layer)
  - w's depend on z's (a hidden layer)
  - $ightharpoonup \mathbf{z}'s$  depend on  $\mathbf{x}'s$ .
- Neural nets are good for prediction
  - especially in speech recognition (Google Translate), image recognition,
  - but require much tuning and very difficult (impossible) to interpret
  - and basis for deep nets and deep learning.



## Neural Network Example



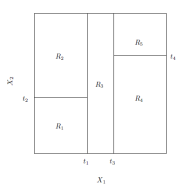
hidden layer 1 hidden layer 2

## 5. Regression Trees and Random Forests

- Regression trees sequentially split regressors x into regions that best predict y.
- ullet Sequentially split  ${f x}'s$  into rectangular regions in way that reduces RSS
  - then  $\hat{y}_i$  is the average of y's in the region that  $\mathbf{x}_i$  falls in
  - with J blocks RSS=  $\sum_{j=1}^{J} \sum_{i \in R_j} (y_i \bar{y}_{R_j})^2$ .
- Simplest case is a single x
  - split at  $x^*$  that minimizes  $\sum_{i:x_i \leq x^*} (y_i \bar{y}_{R_1})^2 + \sum_{i:x_i > x^*} (y_i \bar{y}_{R_1})^2$ 
    - ★ where  $\bar{y}_{R_1}$  is average of  $y_i$  for  $i: x_i \leq x^*$
    - ★ and  $\bar{y}_{R_2}$  is average of  $y_i$  for  $i: x_i > x^*$ .
  - second split is then best split within R<sub>1</sub> and R<sub>2</sub>
  - then predicted y's are a step function of x.

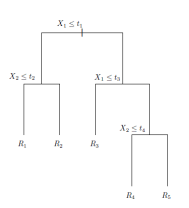
#### Tree example from ISL page 308

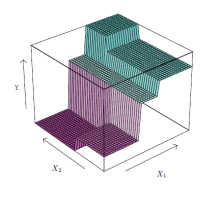
- (1) split X1 in two at  $t_1$ ;
  - (2) split the lowest X1 values into R1 and R2 on basis of X2  $\geq t_2$ ;
  - (3) split the highest X1 values at  $t_3$  into R3 and R4/R5;
  - (4) split the highest X1 values on basis of X2  $\geq t_4$  into R4 and R5.



## Tree example from ISL (continued)

- The left figure gives the tree.
- The right figure shows the predicted values of y.





#### Improvements to regression trees

- Regression trees are easy to understand if there are few regressors.
- But they do not predict as well as methods given so far
  - due to high variance (e.g. split data in two then can get quite different trees).
- Better methods are
  - bagging
    - ★ bootstrap aggregating averages regression trees over many samples
  - random forests
    - averages regression trees over many sub-samples
  - boosting
    - ★ trees build on preceding trees (fit residuals not y).

#### Random Forests

- If we bootstrap resample with replacement (bagging) the B estimates are correlated
  - e.g. if a regressor is important it will appear near the top of the tree in each bootstrap sample.
  - the trees look similar from one resample to the next.
- Random forests get bootstrap resamples (like bagging)
  - but within each bootstrap sample use only a random sample of m < p predictors in deciding each split.
  - usually  $m \simeq \sqrt{p}$
  - this reduces correlation across bootstrap resamples.
  - Susan Athey and coauthors are big on random forests.

#### 6. Other Methods

- Principal components
  - $\triangleright$  reduce from p regressors to M < p linear combinations of regressors
- Basis function models
  - scalar case:  $y_i = \beta_0 + \beta_1 b_1(x_i) + \cdots + \beta_K(x_i) + \varepsilon_i$ 
    - \* where  $b_1(\cdot), ..., b_K(\cdot)$  are basis functions that are fixed and known.
  - global polynomial regression
  - splines: step functions, regression splines, smoothing splines
  - wavelets
  - polynomial is global while the others break range of x into pieces.

### Nonparametric and Semiparametric regression

- Nonparametric regression is the most flexible approach
  - for  $f(\mathbf{x}_0) = \mathsf{E}[y|\mathbf{x} = \mathbf{x}_0]$  borrow from observations near to  $\mathbf{x}_0$
  - k-nearest neighbors and kernel-weighted local regression
  - not practical even for moderate  $p = dim(\mathbf{x})$
  - due to the curse of dimensionality
    - $\star$  e.g. if 10 bins in one dimension need  $10^2$  bins in two dimensions, .....
- Semiparametric models provide some structure to reduce the nonparametric component from many dimensions to fewer dimensions (often one).
  - ▶ partially linear models  $y = f(\mathbf{x}, \mathbf{z}) + u = \mathbf{x}' \boldsymbol{\beta} + g(\mathbf{z}) + u$
  - single-index models  $y = g(\mathbf{x}'\boldsymbol{\beta})$ .
  - generalized additive models  $y = g_1(x_1) + g_2(x_2) + \cdots$



#### 7. Classification

- y's are now categorical e.g. binary.
- Interest lies in predicting y using  $\hat{y}$  (classification)
  - whereas economist typically want  $\widehat{Pr}[y = j | \mathbf{x}]$
  - use number misclassified as loss function (not MSE).
- Some methods choose category with highest  $\widehat{\Pr}[y=j|\mathbf{x}]$ 
  - logit, k-nearest neighbors, discriminant analysis
- Support vector machines skip  $\widehat{\Pr}[y=j|\mathbf{x}]$  and directly get  $\widehat{y}$ 
  - can do better.
- Many ML applications are to classification.

### 8. Unsupervised Learning: cluster analysis

- Challenging area: no y, only x.
- Example is determining several types of individual based on responses to many psychological questions.
- Principal components analysis
  - already presented earlier.
- Clustering Methods
  - k-means clustering.
  - hierarchical clustering.

## ISL Figure 10.5

• Data is  $(x_1.x_2)$  with K=2,3 and 4 clusters identified.

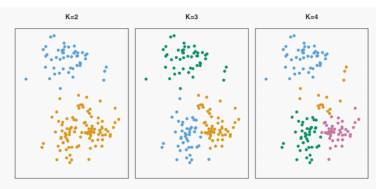


FIGURE 10.5. A simulated data set with 150 observations in two-dimensional space. Panels show the results of applying K-means clustering with different values of K, the number of clusters. The color of each observation indicates the cluster to which it was assigned using the K-means clustering algorithm. Note that

#### 9. ML for Prediction for Economics

- Microeconometrics focuses on estimation of  $\beta$  or of partial effects.
- ullet But in some cases we are directly interested in predicting y
  - probability of one-year survival following hip transplant operation
    - ★ if low then do not have the operation.
  - probability of re-offending
    - ★ if low then grant parole to prisoner.
- Sendhil Mullainathan and J. Spiess: "Machine Learning: An Applied Econometric Approach", Journal of Economic Perspectives, Spring 2017, 87-106.
  - good article to read
  - consider prediction of housing prices
  - detail how to do this using machine learning methods
  - ▶ and then summarize many recent economics ML applications.

### 10. ML for causal effects in partial linear model

- Microeconometrics focuses on estimation of  $\beta$  or of partial effects.
- Consider partial linear model  $y = \beta x_1 + g(\mathbf{x}_2) + u$ .
  - ▶ a good choice of controls  $g(\mathbf{x}_2)$  makes the assumption that  $Cov(x_1, u) = 0$  more plausible
  - so can give  $\beta$  a causal interpretation.
- So use machine learner to come up with good  $g(\mathbf{x}_2)$ 
  - but use ML in a way that akkows valid inference even though we are data mining.
- Belloni, Chernozhukov and Hansen (2014), "High-dimensional methods and inference on structural and treatment effects," *Journal* of Economic Perspectives, Spring, 29-50
  - provides three examples including an IV example.

#### ML for causal treatment effects

- ullet Consider a binary treatment, so  $x_1=d\in\{0,1\}$
- The preceding partially linear model  $y = \beta d + \mathbf{x}_2' \delta + u$ 
  - ightharpoonup restricts the same response  $\beta$  for each individual
  - requires that  $E[u|d, \mathbf{x}_2] = 0$  for unconfoundedness.
- The heterogeneous effects approach is more flexible
  - different responses for different individuals
  - ▶ and unconfoundness assumptions may be more reasonable.
- ML methods have been developed for estimation of the average treatment effect.

#### 11. References

- My website has various detailed slides on machine learning
  - http://cameron.econ.ucdavis.edu/e240f/machinelearning.html
- Chapter 28 of A. Colin Cameron and Pravin K. Trivedi, Microeconometrics using Stata: Volume 2, Stata Press
  - covers machine learning methods for prediction and for causal inference
  - Stata provides a good introduction to machine learning
    - ★ though more advanced ML prediction methods use Python or R.
  - ▶ see https://cameron.econ.ucdavis.edu/mus2/ for book information.
- Standard texts available free at https://www.statlearning.com/ are
  - ► Gareth James, Daniela Witten, Trevor Hastie and Robert Tibshirani (2021), An Introduction to Statistical Learning: with Applications in R, 2nd edition, Springer.
  - ► Gareth James, Daniela Witten, Trevor Hastie, Robert Tibshirani and Jonathan Taylor (2021), *An Introduction to Statistical Learning: with Applications in R*, 2nd edition, Springer.