# Machine Learning for Microeconometrics Part 4: More Prediction Methods 

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## Course Outline

- 1. Variable selection and cross validation
- 2. Shrinkage methods
- ridge, lasso, elastic net
- 3. ML for causal inference using lasso
- OLS with many controls, IV with many instruments
- Part 4: Other methods for prediction
- nonparametric regression, principal components, splines
- neural networks
- regression trees, random forests, bagging, boosting
- 5. More ML for causal inference
- ATE with heterogeneous effects and many controls.
- 6. Classification and unsupervised learning
- classification (categorical $y$ ) and unsupervised learning (no y).


## Introduction

- Basics used OLS regression
- though with potentially rich set of regressors with interactions
- and regularized OLS using lasso, ridge and elastic net.
- Now consider various other flexible methods for regress $y$ on $\mathbf{x}$
- some existed before ML became popular
$\star$ nonparametric and semiparametric regression
* principal components
^ basis functions: polynomials, splines, sieves
- some are more recent
$\star$ neural networks
« regression trees and random forests
- in this part consider supervised learning ( $y$ and $\mathbf{x}$ ) for continuous $y$.
- Based on the two books by Hastie and Tibshirani and coauthors.


## Flexible methods

- These slides present many methods
- Which method is best (or close to best) varies with the application
- e.g. deep learning (neural nets) works very well for Google Translate
- all require setting tuning parameters which may not be straightforward.
- In forecasting competitions the best forecasts are ensembles
- a weighted average of the forecasts obtained by several different methods
- the weights can be obtained by OLS regression in a test sample
$\star$ e.g. given three forecast methods minimize w.r.t. $\tau_{1}$ and $\tau_{2}$

$$
\sum_{i=1}^{n}\left\{y_{i}-\tau_{1} \widehat{y}_{i}^{(1)}-\tau_{2} \widehat{y}_{i}^{(2)}-\left(1-\tau_{1}-\tau_{2}\right) \widehat{y}_{i}^{(3)}\right\}^{2} .
$$

## Implementation

- Stata has built-in commands for
- lasso and ridge (already covered)
- nonparametric regression, principal components and basis functions.
- For other models (neural network, random forests, ...)
- R has many commands
- Python is viewed as best
- Stata has add-ons that often are front-ends to R or Python so require also installing R or Python
* crtrees does trees and random forests directly in Stata
* srtree is a wrapper for R functions tree(), randomForest(), and gbm()
* pylearn does trees, random forests and neural nets directly in Stata and requires installation of Python and the Python scikit-learn library.


## Overview

(1) Nonparametric and semiparametric regression
(2) Dimension reduction (principal components)
(3) Flexible regression (polynomials, splines, sieves)
(1) Neural networks
(5) Regression trees and random forests
(1) Regression trees
(2) Bagging
© Random forests

- Boosting
(0) Prediction Example
(3) Prediction for Economics


## 1. Nonparametric and Semiparametric Regression

- We want a flexible model for $f\left(\mathbf{x}_{0}\right)=E\left[y \mid \mathbf{x}=\mathbf{x}_{0}\right]$ evaluated at a range of values $\mathbf{x}_{0}$.
- Use $\bar{y} \mid \mathbf{x}=\mathbf{x}_{0}$ if there are many observations of $y$ at each value of $\mathbf{x}_{0}$.
- In practice there are few values of $y$ at each value of $\mathbf{x}_{0}$,
so bring into the average values of $y$ at values of $x$ near to $x_{0}$
- local constant kernel-weighted regression
- local linear kernel-weighted regression
- $k$-nearest neighbors
- lowess.
- All depend on a tuning parameter that trades off bias and variance
- like choice of bin width for a histogram
- usually minimize MSE using leave-one-out cross validation.
- Problem: curse of dimensionality if many $\mathbf{x}$
- Solutions: semiparametric (old school) or ML methods (new school).


### 1.1 Nonparametric kernel-weighted regression

- Start with scalar $x$.
- Local-weighted average

$$
\widehat{f}\left(x_{0}\right)=\sum_{i=1}^{N} w\left(x_{i}, x_{0}, h\right) y_{i}, \sum_{i=1}^{N} w_{i}=1
$$

- Kernel-weighted average uses kernel weights

$$
w\left(x_{i}, x_{0}, h\right)=K\left(\frac{x_{i}-x_{0}}{h}\right) /\left(\sum_{j=1}^{N} K\left(\frac{x_{j}-x_{0}}{h}\right)\right)
$$

- $K(z)$ is a kernel function with $\int K(z) d z=1, K(-z)=K(z)$, $K(z) \rightarrow 0$ as $z \rightarrow \infty$.
- Most kernels have $K(z)=0$ for $|z|>1$ (epanechnikov, uniform, triangular)
- Others are continuous on $(-\infty, \infty)$ (gaussian).
- Key is the bandwidth $h$ (the tuning parameter)
- chosen by leave-one-out cross validation (minimizes MSE).


## Local constant regression

- Recall: a sample mean of $y=$ OLS of $y$ on an intercept.
- Similarly: a weighted sample mean of $y=$ weighted OLS of $y$ on an intercept.
- Weighted regression on a constant is the estimator that minimizes the weighted sum of squared residuals

$$
\sum_{i=1}^{N} w\left(x_{i}, x_{0}, h\right)\left(y_{i}-\alpha_{0}\right)^{2}
$$

- which yields $\widehat{\alpha}_{0}=\sum_{i=1}^{N} w\left(x_{i}, x_{0}, h\right) y_{i}$ which is the estimator in the previous slide.
- so called (kernel-weighted) local constant regression.


## Local linear regression

- The local linear estimator generalizes to linear regression in the neighborhood of $x_{0}$.
- Then $\widehat{f}\left(x_{0}\right)=\widehat{\alpha}_{0}$ where minimize w.r.t. $\alpha_{0}$ and $\beta_{0}$

$$
\sum_{i=1}^{N} w\left(x_{i}, x_{0}, h\right)\left\{y_{i}-\alpha_{0}-\beta_{0}\left(x_{i}-x_{0}\right)\right\}^{2} .
$$

- so called (kernel-weighted) local linear regression.
- Advantages
- better estimates at endpoints of the data
- $\widehat{\beta}_{0}=\widehat{f}^{\prime}\left(x_{0}\right)$ provides an estimate of the gradient $\partial E[y \mid x] /\left.\partial x\right|_{x_{0}}$.
- Stata commands
- lpoly y x, degree(1)
- or npregress kernel y x, estimator(linear)


## Other nonparametric methods

- Lowess (locally weighted scatterplot smoothing)
- a variation of local linear with variable bandwidth, tricubic kernel and downweighting of outliers
- lowess y x.
- k-nearest neighbors
- use an equal-weighted average of $y$ values for the $k$ observations with $x_{i}$ closest to $x_{0}$
- $\widehat{f}\left(\mathbf{x}_{0}\right)=\frac{1}{k} \sum_{i=1}^{N} \mathbf{1}\left[\mathbf{x}_{i} \in N\left(\mathbf{x}_{0}\right)\right] \times y_{i}$
- Stata user-written command knnreg
- not used much in economics aside from matching estimates of ATE.


### 1.2 Curse of Dimensionality

- Nonparametric methods do not extend well to multiple regressors.
- Consider p-dimensional $\mathbf{x}$ broken into bins
- for $p=1$ we might average $y$ in each of 10 bins of $x$
- for $p=2$ we may need to average over $10^{2}$ bins of $\left(x_{1}, x_{2}\right)$
- and so on.
- On average there may be few to no points with high-dimensional $\mathbf{x}_{i}$ close to $\mathrm{x}_{0}$
- called the curse of dimensionality.
- Formally for local constant kernel regression with bandwidth $h$
- bias is $O\left(h^{2}\right)$ and variance is $O\left(n h^{p}\right)$
- optimal bandwidth is $O\left(n^{-1 /(p+4)}\right)$
$\star$ gives asymptotic bias so standard conf. intervals not properly centered
- convergence rate is then $n^{-2 /(p+4)} \ll n^{-0.5}$


### 1.3 Kernel Regression in Higher Dimensions

- Kernel regression extends to higher dimensions.
- For $\operatorname{dim}(\mathbf{x})=k$ local linear regression $\alpha_{0}$ and $\beta_{0}$ minimize

$$
\sum_{i=1}^{N} w\left(\mathbf{x}_{i}, \mathbf{x}_{0}, \mathbf{h}\right)\left\{y_{i}-\alpha_{0}-\left(\mathbf{x}_{i}-\mathbf{x}_{0}\right)^{\prime} \boldsymbol{\beta}_{0}\right\}^{2}
$$

where $w\left(\mathbf{x}_{i}, \mathbf{x}_{0}, \mathbf{h}\right)=\prod_{j=1}^{k} w\left(x_{j i}, x_{j 0}, h_{j}\right)$ is a product kernel.

- Then $\widehat{\alpha}_{0}=\widehat{f}\left(\mathbf{x}_{0}\right)$ can predict $f\left(\mathbf{x}_{0}\right)$ poorly due to curse of dimensionality.
- But when we average we may predict better.
- In particular can compute quantities such as
- average marginal effect for the $j^{t h}$ regressor
- marginal effect for the $j^{\text {th }}$ regressor with other regressors set at their mean values or a prespecified value
- Stata's npregress kernel command does this
- use bootstrap to get confidence intervals.


### 1.4 Semiparametric Models

- Semiparametric methods place some structure on the problem
- parametric component $(\boldsymbol{\beta})$ for part of the model
- nonparametric component (function(s) $f$ ) that is often one dimensional
- Ideally $\sqrt{N}(\widehat{\boldsymbol{\beta}}-\boldsymbol{\beta}) \xrightarrow{d} \mathcal{N}[\mathbf{0}, \mathbf{V}]$ despite the nonparametric component.
- Leading examples
- partial linear (used in economics)
- single-index (used in economics)
- generalized additive model (used in statistics)
- project pursuit (used in statistics).


## Leading semiparametric models

- Partial linear model: $E\left[y_{i} \mid \mathbf{x}_{i}, \mathbf{z}_{i}\right]=\mathbf{x}_{i}^{\prime} \boldsymbol{\beta}+g\left(\mathbf{z}_{i}\right)$ where $g(\cdot)$ not specified.
- use Robinson differencing estimator.
- Single index model: $\mathrm{E}\left[y_{i} \mid \mathbf{x}_{i}\right]=g\left(\mathbf{x}_{i}^{\prime} \boldsymbol{\beta}\right)$ where $g(\cdot)$ not specified
- Ichimura semiparametric least squares
$\star \widehat{\boldsymbol{\beta}}$ and $\widehat{g}$ minimize $\sum_{i=1}^{N} w\left(\mathbf{x}_{i}\right)\left\{y_{i}-\widehat{g}\left(\mathbf{x}_{i}^{\prime} \boldsymbol{\beta}\right)\right\}^{2}$
$\star$ where $w\left(\mathbf{x}_{i}\right)$ is a trimming function that drops outlying $\mathbf{x}$ values.
- can only estimate $\beta$ up to scale in this model
* still useful as ratio of coefficients equals ratio of marginal effects in a single-index models.
- Generalized additive model: $E\left[y_{i} \mid \mathbf{x}_{i}\right]=g_{1}\left(x_{1 i}\right)+\cdots+g_{K}\left(x_{K i}\right)$ where the $g_{j}(\cdot)$ are unspecified.
- estimate by backfitting
- can make more complex by e.g. $x_{3 i}=x_{1 i} \times x_{2 i}$.


### 1.5 How can ML methods do better?

- Machine learning methods can outperform nonparametric and semiparametric methods
- so wherever econometricians use nonparametric and semiparametric regression in higher-dimensional models it may be useful to use ML methods.
- In theory there is scope for improving nonparametric methods.
- k-nearest neighbors usually has a fixed number of neighbors
- but it may be better to vary the number of neighbors with data sparsity
- Kernel-weighted local regression methods usually use a fixed bandwidth
- but it may be better to vary the bandwidth with data sparsity.
- There may be advantage to basing neighbors in part on relationship with $y$.


## 2. Dimension Reduction

- Reduce from $p$ regressors to $M<p$ linear combinations of regressors.
- So form

$$
\underset{(N \times m)}{\mathbf{X}^{*}}=\underset{(N \times p)}{\mathbf{X}} \times \underset{(p \times M)}{\mathbf{A}} \text { where } M<p .
$$

- Then after dimension reduction

$$
\begin{aligned}
\mathbf{y} & =\beta_{0}+\mathbf{X}^{*} \delta+\mathbf{u} \\
& =\beta_{0}+\mathbf{X} \boldsymbol{\beta}+\mathbf{u} \text { where } \boldsymbol{\beta}=\mathbf{A} \boldsymbol{\delta} .
\end{aligned}
$$

- Aside: in ML high-dimensional simply means $p$ is large relative to $n$
- some methods (not PCA) even allow $p>n$
- $n$ could be large or small.


## Dimension Reduction (continued)

- Two methods are covered in ISL
- 1. Principal components (PCA)
$\star$ use only $\mathbf{X}$ to form $\mathbf{A}$ (unsupervised)
- 2. Partial least squares (PLS)
$\star$ also use relationship between $\mathbf{y}$ and $\mathbf{X}$ to form $\mathbf{A}$ (supervised).
- PCA is widely used in other social sciences
- and is related to factor analysis.
- PCA can work remarkably well even though it ignores $y$
- intuitively partial LS should be better
- but in practice not so as partial LS can add variance
- I have not seen partial LS used in practice - can skip.
- For PCA and PLS standardize regressors as not scale invariant
- and can use cross-validation to determine $M$.


### 2.1 Principal Components Analysis (PCA)

- Suppose $\mathbf{X}$ is normalized to have zero means so $i j^{t h}$ entry is $x_{j i}-\bar{x}_{j}$.
- The first principal component has the largest sample variance among all normalized linear combinations of the columns of $n \times p$ matrix $\mathbf{X}$
- the first component is $\mathbf{X h}_{1}$ where $\mathbf{h}_{1}$ is $p \times 1$
- normalize $\mathbf{h}_{1}$ so that $\mathbf{h}_{1}^{\prime} \mathbf{h}_{1}=1$
- then $\mathbf{h}_{1}$ max $\operatorname{Var}\left(\mathbf{X} \mathbf{h}_{1}\right)=\mathbf{h}_{1}^{\prime} \mathbf{X}^{\prime} \mathbf{X} \mathbf{h}_{1}$ subject to $\mathbf{h}_{1}^{\prime} \mathbf{h}_{1}=1$
- the maximum is the largest eigenvalue of $\mathbf{X}^{\prime} \mathbf{X}$ and $\mathbf{h}_{1}$ is the corresponding eigenvector.
- The second principal component has the largest variance subject to being orthogonal to the first, and so on.


## Formulas for PCA

- Eigenvalues and eigenvectors of $\mathbf{X}^{\prime} \mathbf{X}$
- Let $\Lambda=\operatorname{Diag}\left[\lambda_{j}\right]$ be $p \times p$ vector of eigenvalues of $\mathbf{X}^{\prime} \mathbf{X}$
- Order so $\lambda_{1} \geq \lambda_{2} \geq \cdots \geq \lambda_{1}$
- Let $\mathbf{H}=\left[\mathbf{h}_{1} \cdots \mathbf{h}_{p}\right]$ be $p \times p$ vector of corresponding eigenvectors
- $\mathbf{X}^{\prime} \mathbf{X} \mathbf{h}_{1}=\lambda_{1} \mathbf{h}_{1}$ and $\mathbf{X}^{\prime} \mathbf{X H}=\Lambda \mathbf{H}$ and $\mathbf{H}^{\prime} \mathbf{H}$
- Then
- the $j^{\text {th }}$ principal component is $\mathbf{X h}_{j}$
- $M$-principal components regression uses $\mathbf{X}^{*}=\mathbf{X A}$ where $\mathbf{A}=\left[h_{1} \cdots h_{M}\right]$.
- ASIDE: PCA is related to, but not exactly the same as factor analysis
- factor analysis decomposes observed $p$-dimensional $\mathbf{x}$ into a linear combination of $M$ unobserved factor loadings and $p$ i.i.d. errors
- often $M=1$ so $x_{j i}=$ a multiple of the one common factor plus i.i.d. noise.


## Principal Components Analysis Example

- Stata command pca by default standardizes the data.
- For the d.g.p. for $x_{1}, x_{2}, x_{3}$ (the d.g.p. in part 1 of these slides) we expect eigenvalues $2,0.5$ and 0.5 as $n \longrightarrow \infty$.
. * Principal components with default correlation option that standardizes data
- pca x1 $\times 2 \times 3$

Principa1 components (eigenvectors)

| Variab1e | Comp1 | Comp2 | Comp3 | Unexp1ained |
| ---: | ---: | ---: | ---: | ---: |
| x1 | 0.6306 | -0.1063 | -0.7688 | 0 |
| x2 | 0.5712 | -0.6070 | 0.5525 | 0 |
| x3 | 0.5254 | 0.7876 | 0.3220 | 0 |

## Principal Components Analysis Example (continued)

- First principal component is $0.6306 z x_{1}+0.5712 z x_{2}+0.5254 z x_{3}$
- where $z x_{j}$ are standardized
- and has variance 1.8618 that explains $1.8618 / 3=0.6056$ of the variance.
- The principal components have means 0 , variances equal to the eigenvalue, and are uncorrelated.

```
. * Generate the 3 principal components and their means, st.devs., correlations
. quietly predict pc1 pc2 pc3
. summarize pc1 pc2 pc3
\begin{tabular}{r|rrrrr} 
Variab7e & Obs & Mean & Std. Dev. & Min & Max \\
\hline pc1 & 40 & \(-3.35 \mathrm{e}-09\) & 1.347842 & -2.52927 & 2.925341 \\
pc2 & 40 & \(-3.63 \mathrm{e}-09\) & .8529281 & -1.854475 & 1.98207 \\
pc3 & 40 & \(2.08 \mathrm{e}-09\) & .6751564 & -1.504279 & 1.520466
\end{tabular}
. correlate pc1 pc2 pc3
(obs=40)
\begin{tabular}{r|rrr} 
& pc1 & pc2 & pc3 \\
\hline pc1 & 1.0000 & & \\
pc2 & 0.0000 & 1.0000 & \\
pc3 & -0.0000 & -0.0000 & 1.0000
\end{tabular}
```


## Principal Components Analysis Example (continued)

- Correlation coefficient of $y$ with $\widehat{y}$
- $r=0.4871$ from OLS on all three regressors
- $r=0.4444$ from OLS on first principal component
- $r=0.4740$ on just x 1 (the d.g.p. was $y_{i}=2+x_{1 i}+u_{i}$ ).
. * Compare R from OLS on all three regressors, on pc 1 , on x 1 , on x 2 , on x 3 . qui regress y x1 x2 x3
- predict yhat
(option xb assumed; fitted values)
. correlate y yhat pc1 x1 x2 x3
(obs=40)

|  | $y$ | $y h a t$ | $p c 1$ | $x 1$ | $x 2$ | $x 3$ |
| ---: | ---: | ---: | ---: | ---: | ---: | ---: |
| $y$ | 1.0000 |  |  |  |  |  |
| yhat | 0.4871 | 1.0000 |  |  |  |  |
| pc1 | 0.4444 | 0.9122 | 1.0000 |  |  |  |
| x1 | 0.4740 | 0.9732 | 0.8499 | 1.0000 |  |  |
| x2 | 0.3370 | 0.6919 | 0.7700 | 0.5077 | 1.0000 |  |
| x3 | 0.2046 | 0.4200 | 0.7082 | 0.4281 | 0.2786 | 1.0000 |

## Principal Components Analysis (continued)

- PCA is unsupervised so seems unrelated to $\mathbf{y}$ but
- Elements of Statistical Learning says does well in practice.
- PCA has the smallest variance of any estimator that estimates the model $\mathbf{y}=\mathbf{X} \boldsymbol{\beta}+\mathbf{u}$ with i.i.d. errors subject to constraint $\mathbf{C} \boldsymbol{\beta}=\mathbf{c}$ where $\operatorname{dim}[\mathbf{C}] \leq \operatorname{dim}[\mathbf{X}]$.
- PCA discards the $p-M$ smallest eigenvalue components whereas ridge does not, though ridge does shrink towards zero the most for the smallest eigenvalue components (ESL p.79).
- For machine learning the tuning parameter is the number of components and use e.g. K-fold class validation to determine this.
- For completeness next give partial least squares which is supervised.


### 2.2 Partial Least Squares

- Partial least squares produces a sequence of orthogonal linear combinations of the regressors.
- 1. Standardize each regressor to have mean 0 and variance 1 .
- 2. Regress $y$ individually on each $\mathbf{x}_{j}$ and let $\mathbf{z}_{1}=\sum_{j=1}^{p} \hat{\theta}_{1 j} \mathbf{x}_{j}$
- 3. Regress $y$ on $\mathbf{z}_{1}$ and let $\widehat{\mathbf{y}}^{(1)}$ be prediction of $\mathbf{y}$.
- 4. Orthogonalize each $\mathbf{x}_{j}$ by regress on $\mathbf{z}_{1}$ to give $\mathbf{x}_{j}^{(1)}=\mathbf{x}_{j}-\mathbf{z}_{1} \widehat{\tau}_{j}$ where $\widehat{\tau}_{j}=\left(\mathbf{z}_{1}^{\prime} \mathbf{z}_{1}\right)^{-1} \mathbf{z}_{1}^{\prime} \mathbf{x}_{j}^{(1)}$.
- 5. Go back to step 1 with $\mathbf{x}_{j}$ now $\mathbf{x}_{j}^{(1)}$, etc.
- When done $\widehat{\mathbf{y}}=\widehat{\mathbf{y}}^{(1)}+\widehat{\mathbf{y}}^{(2)}+\cdots$
- Partial least squares turns out to be similar to PCA
- especially if $R^{2}$ is low.


## 3. Flexible Regression using Basis Functions

- Basis function models (or sieves)
- global polynomial regression
- splines: step functions, regression splines, smoothing splines, b-splines
- polynomial is global while the others break range of $x$ into pieces.
- Can make nonparametric
- increase order of polynomial or number of knots (split points) in splines
- select model using leave-one-out cross validation, generalized cross validation, Mallows CP, AIC or BIC
- Stata npregress series command implements these methods
- options polynomial, spline, bspline
- nonparametric tuning use option criterion()
- or parametric tuning use options polynomial(\#), spline(\#), knots(\#).


### 3.1 Basis Functions

- Also called series expansions and sieves.
- General approach (scalar $x$ for simplicity)

$$
y_{i}=\beta_{0}+\beta_{1} b_{1}\left(x_{i}\right)+\cdots+\beta_{K} b_{K}\left(x_{i}\right)+\varepsilon_{i}
$$

- where $b_{1}(\cdot), \ldots, b_{K}(\cdot)$ are basis functions that are fixed and known.
- Global polynomial regression sets $b_{j}\left(x_{i}\right)=x_{i}^{j}$
- typically $K \leq 3$ or $K \leq 4$.
- fits globally and can overfit at boundaries.
- Step functions: separately fit $y$ in each interval $x \in\left(c_{j}, c_{j+1}\right)$
- could be piecewise constant or piecewise linear.
- Splines smooth so that not discontinuous at the cut points.
- Wavelets are also basis functions, richer than Fourier series.


## Global Polynomials Example

- Generated data:

$$
y_{i}=1+x_{1 i}+x_{2 i}+f\left(z_{i}\right)+u_{i} \text { where } f(z)=z+z^{2} .
$$

. * Generated data: $y=1+1 * x 1+1 * x 2+f(z)+u$ where $f(z)=z+z \wedge 2$
. clear
. set obs 200
number of observations (_N) was 0, now 200
. set seed 10101

- generate $x 1=$ rnormal ()
. generate $\times 2=$ rnormal ()$+0.5 * x 1$
. generate $z=\operatorname{rnormal}()+0.5 * x 1$
. generate $z s q=z \wedge 2$
. generate $y=1+x 1+x 2+z+z s q+2$ *rnormal ()
. summarize

| Variable | Obs | Mean | Std. Dev. | Min | Max |
| ---: | ---: | ---: | ---: | ---: | ---: |
| x1 | 200 | .0301211 | 1.014172 | -3.170636 | 3.093716 |
| x2 | 200 | .0226274 | 1.158216 | -4.001105 | 3.049917 |
| z | 200 | .0664539 | 1.146429 | -3.386704 | 2.77135 |
| zsq | 200 | 1.312145 | 1.658477 | .0000183 | 11.46977 |
| $y$ | 200 | 2.164401 | 3.604061 | -5.468721 | 14.83116 |

## Global Polynomials Example (continued)

- Fit quartic in $z$ (with $x_{1}$ and $x_{2}$ omitted) and compare to quadratic in $z$
- regress y c.z\#\#c.z\#\#c.z\#\#c.z, vce(robust)
- quartic chases endpoints.



### 3.2 Regression Splines

- Begin with step functions: separate fits in each interval $\left(c_{j}, c_{j+1}\right)$
- Piecewise constant
- $b_{j}\left(x_{i}\right)=1\left[c_{j} \leq x_{i}<c_{j+1}\right]$
- Piecewise linear
- intercept is $1\left[c_{j} \leq x_{i}<c_{j+1}\right]$ and slope is $x_{i} \times 1\left[c_{j} \leq x_{i}<c_{j+1}\right]$
- Problem is that discontinuous at the cut points (does not connect)
- solution is splines.


## Piecewise linear spline

- Begin with piecewise linear with two knots at $c$ and $d$

$$
\begin{aligned}
f(x)= & \alpha_{1} 1[x<c]+\alpha_{2} 1[x<c] x+\alpha_{3} 1[c \leq x<d] \\
& +\alpha_{4} 1[c \leq x<d] x+\alpha_{5} 1[x \geq d]+\alpha_{6} 1[x \geq d] x .
\end{aligned}
$$

- To make continuous at $c$ (so $f(c-)=f(c))$ and $d$ (so $f(d-)=f(d))$ we need two constraints

$$
\begin{array}{lr}
\text { at } c: & \alpha_{1}+\alpha_{2} c=\alpha_{3}+\alpha_{4} c \\
\text { at } d: & \alpha_{3}+\alpha_{4} d=\alpha_{5}+\alpha_{6} d .
\end{array}
$$

- Alternatively introduce the Heaviside step function

$$
h_{+}(x)=x_{+}= \begin{cases}x & x>0 \\ 0 & \text { otherwise }\end{cases}
$$

- Then the following imposes the two constraints (so have 6-2 $=4$ regressors)

$$
f(x)=\beta_{0}+\beta_{1} x+\beta_{2}(x-c)_{+}+\beta_{2}(x-d)_{+}
$$

## Spline Example

- Piecewise linear spline with two knots done manually.


```
* Create the basis function manually with three segments and knots at -1 and 1
generate zsegl = z
. replace zseg2 = z - (-1) if z > -1
163 real changes made)
. generate zseg3 = 0
rep1ace zseg3 = z - 1 if z > 1
(47 real changes made)
```

. * Piecewise linear regression with three sections
regress y zseg1 zseg2 zseg3

## Spline Example (continued)

- Plot of fitted values from piecewise linear spline has three connected line segments.



## Spline Example (continued)

- The mkspline command creates the same spline variables.
. * Repeat piecewise linear using command mkspline to create the basis functions
. mksp1ine zmk1 -1 zmk2 1 zmk3 = z, margina1
. summarize zseg1 zmk1 zseg2 zmk2 zseg3 zmk3, sep (8)

| Variable | Obs | Mean | Std. Dev. | Min | Max |
| ---: | ---: | ---: | ---: | ---: | ---: |
| zseg1 | 200 | .0664539 | 1.146429 | -3.386704 | 2.77135 |
| zmk1 | 200 | .0664539 | 1.146429 | -3.386704 | 2.77135 |
| zseg2 | 200 | 1.171111 | .984493 | 0 | 3.77135 |
| zmk2 | 200 | 1.171111 | .984493 | 0 | 3.77135 |
| zseg3 | 200 | .138441 | .3169973 | 0 | 1.77135 |
| zmk3 | 200 | .138441 | .3169973 | 0 | 1.77135 |

- To repeat earlier results: regress y zmk1 zmk2 zmk3
- And to add regressors: regress y x1 x2 zmk1 zmk2 zmk3


## Cubic Regression Splines

- This is the standard.
- Piecewise cubic spline with $K$ knots
- require $f(x), f^{\prime}(x)$ and $f^{\prime \prime}(x)$ to be continuous at the $K$ knots
- Then can do OLS with

$$
f(x)=\beta_{0}+\beta_{1} x+\beta_{2} x^{2}+\beta_{3} x^{3}+\beta_{4}\left(x-c_{1}\right)_{+}^{3}+\cdots+\beta_{(3+K)}\left(x-c_{K}\right)_{+}^{3}
$$

- for proof when $K=1$ see ISL exercise 7.1.
- This is the lowest degree regression spline where the graph of $\widehat{f}(x)$ on $x$ seems smooth and continuous to the naked eye.
- There is no real benefit to a higher-order spline.
- Regression splines overfit at boundaries.
- A natural (or restricted) cubic spline is an adaptation that restricts the relationship to be linear past the lower and upper boundaries of the data.


## Spline Example

- Natural or restricted cubic spline with five knots at the 5, 27.5,50, 72.5 and 95 percentiles
- mkspline zspline = z, cubic nknots(5) displayknots
- regress y zspline*



## Other Splines

- Regression splines and natural splines require choosing the cut points
- e.g. use quintiles of $x$.
- Smoothing splines avoid this
- use all distinct values of $x$ as knots
- but then add a smoothness penalty that penalizes curvature.
- The function $g(\cdot)$ minimizes

$$
\sum_{i=1}^{n}\left(y_{i}-g\left(\mathbf{x}_{i}\right)\right)^{2}+\lambda \int_{a}^{b} g^{\prime \prime}(t) d t \text { where } a \leq \text { all } x_{i} \leq b
$$

- $\lambda=0$ connects the data points and $\lambda \rightarrow \infty$ gives OLS.
- For multivariate splines use multivariate adaptive regression splines (MARS).


## Stata Commands

- The preceding examples were done manually for pedagogical reasons.
- Stata's npregress series command has options
- polynomial(\#) for a global polynomial of order \#
- spline(\#) for a natural spline of order \#
- bspline(\#) for a b-spline of order \#
- For splines and B-splines the number of knots can be determined
- by CV (the default), generalized CV, AIC, BIC or Cp
- option knots(\#) where \# is the number of knots
- option knotsmat (matname) specifies the values of the knots.
- If there is more than one regressor then the basis functions for each regressor may be interacted or not interacted.
- Stata user-written add-on commands
- gam (Royston and Ambler) for smoothing splines
- bspline command (Newson 2012) for a range of bases.


### 3.3 Wavelets

- Wavelets are used especially for signal processing and extraction.
- they are richer than a Fourier series basis.
- they can handle both smooth sections and bumpy sections of a series.
- they are not used in cross-section econometrics
- they may be useful for some time series.
- Start with a mother or father wavelet function $\psi(x)$.
- example is the Haar function $\psi(x)=\left\{\begin{array}{cc}1 & 0 \leq x<\frac{1}{2} \\ -1 & \frac{1}{2}<x<1 \\ 0 & \text { otherwise }\end{array}\right.$
- Then both translate by $b$ and scale by $a$ to give basis functions $\psi^{a b}(x)=|a|^{-1 / 2} \psi\left(\frac{x-b}{a}\right)$.


## 4. Neural Networks and Deep Learning

- Deep learning is learning that occurs in a series of levels or layers that can go to considerable depth.
- The leading example is a neural network with multiple hidden layers.
- The term neural arises as initial models were based on mimicking how the brain works.
- Neural networks have been around for a long time
- but to work well they need to be complex with many parameters
- this requires a lot of data and good computational techniques.
- Their use has exploded in the past fifteen years
- due to more computational power, more data, better algorithms, newer models
- they work especially well for image recognition and language translation (Google Translate).


### 4.1 Hidden Layer(s) Neural Networks

- A neural network involves a series of nested regression models.
- A single hidden layer neural network explaining $y$ by $\mathbf{x}$ has
- $y$ depends on $z^{\prime} s$ (hidden units)
- $\mathbf{z}^{\prime} s$ depend on $\mathbf{x}^{\prime} s$ (input units).
- A neural network with two hidden layers explaining $y$ by $\mathbf{x}$ has
- y depends on $\mathbf{w}$ 's (hidden units)
- w's depend on $z^{\prime} s$ (hidden units)
- $z^{\prime} s$ depend on $x^{\prime} s$ (input units).


## Two hidden layer neural network

Input $\mathbf{x} \rightarrow \mathbf{z} \rightarrow \mathbf{w} \rightarrow y$ output.


## One hidden layer neural network

- $y$ depends on $M \mathbf{z}^{\prime} s$ and the $\mathbf{z}^{\prime} s$ depend on $p \mathbf{x}^{\prime} s(y \leftarrow \mathbf{z} \leftarrow \mathbf{x})$.
- Then

$$
\begin{aligned}
f(\mathbf{x}) & =\beta_{0}+\mathbf{z}^{\prime} \boldsymbol{\beta}=\beta_{0}+\sum_{m=1}^{M} \beta_{m} z_{m}(\mathbf{x}) \\
z_{m}(\mathbf{x}) & =g\left(\alpha_{0 m}+\mathbf{x}^{\prime} \boldsymbol{\alpha}_{m}\right)=g\left(\alpha_{0 m}+\sum_{j=1}^{p} \alpha_{m j} x_{j}\right), m=1, \ldots, M
\end{aligned}
$$

- The $z_{m}(\mathbf{x})$ functions are called activations.
- The specified nonlinear function $g(\cdot)$ is called an activation function.
- Initially the sigmoid activation function was used

$$
g(v)=\frac{1}{1+e^{-v}} .
$$

- Now use the rectified linear unit (ReLU) activation function

$$
g(v)=\max (0, v)
$$

### 4.2 Computation

- A neural network model has a highly nonlinear and nonconvex objective function
- Using squared error loss we minimize $\sum_{i=1}^{n}\left(y_{i}-f\left(\mathbf{x}_{i}, \boldsymbol{\theta}\right)\right)^{2}$.
- With one hidden layer $\beta_{0}, \ldots, \beta_{m}, \alpha_{10}, \ldots, \alpha_{1 p}, \ldots, \alpha_{M 0}, \ldots, \alpha_{M p}$ minimize

$$
Q(\boldsymbol{\theta})=\sum_{i=1}^{n}\left\{y_{i}-\beta_{0}-\sum_{m=1}^{M} \beta_{m}\left[g\left(\alpha_{m 0}+\sum_{j=1}^{p} \alpha_{m j} x_{j}\right)\right]\right\}^{2} .
$$

- Optimization uses back propagation stochastic gradient descent.
- Back propagation computes gradients using the chain rule

$$
\frac{\partial Q_{i}(\boldsymbol{\theta})}{\partial \alpha_{m j}}=-2\left(y_{i}-f\left(\mathbf{x}_{i}\right)\right) \times \boldsymbol{\beta}_{m} \times \frac{\partial g\left(\mathbf{x}_{i}^{\prime} \boldsymbol{\alpha}\right)}{\partial \alpha_{m j}} \times x_{i j}
$$

- This saves computation time as e.g. $-2\left(y_{i}-f\left(\mathbf{x}_{i}\right)\right) \times \boldsymbol{\beta}_{m}$ is the same for all $\alpha_{m 1}, \ldots, \alpha_{m p}$.
- "Forward propagation" goes from inputs to output: $x \rightarrow z \rightarrow \hat{y}$
- Called back propagation as gradient computation order is $\hat{y} \rightarrow z \rightarrow x$.


## Computation (continued)

- Stochastic gradient descent uses update rule

$$
\widehat{\boldsymbol{\theta}}_{s+1}=\widehat{\boldsymbol{\theta}}_{s}-\left.\lambda_{s} \frac{\partial Q_{s}(\boldsymbol{\theta})}{\partial(\boldsymbol{\theta})}\right|_{\hat{\boldsymbol{\theta}}_{s}}
$$

- Key is that the gradient is computed at each round using only a small subsample (that varies across each round $s$ )
- $Q_{s}(\cdot)$ is a small randomly-chosen subsample of the data
- we don't have to exactly estimate the gradient at each step
- though then have many iterations.
- Stopping rule is e.g. cross validation measure is minimized
- rather than the usual gradient is zero.
- To avoid overfitting may use a Ridge or Lasso penalty in $Q(\boldsymbol{\theta})$.
- In econometrics we instead use $\widehat{\boldsymbol{\theta}}_{s+1}=\widehat{\boldsymbol{\theta}}_{s}-H_{s}^{-1} \times\left.\frac{\partial Q(\boldsymbol{\theta})}{\partial(\boldsymbol{\theta})}\right|_{\hat{\boldsymbol{\theta}}_{s}}$
- computationally expensive for high-dimensional $\boldsymbol{\theta}$
- and $H_{s}$ can be ill-conditioned (multicollinearity).


## Computation (continued)

- Neural nets require a lot of fine tuning - not off-the-shelf.
- We need to determine
- the number of hidden layers
- the number $M$ of hidden units within each layer
* best to have too many and then avoid overfit using penalty.
- have a good algorithm to compute $\alpha^{\prime} s, \beta^{\prime} s, \ldots$.
$\star$ the subsample size for stochastic gradient descent
$\star$ the step size $\lambda_{s}$ (which should decrease with $s$ )
$\star$ starting values for the $\alpha^{\prime} s, \beta^{\prime} s, \ldots$


### 4.3 Neural Networks Example

- This example uses user-written Stata command brain (Doherr)
- there is one hidden layer with 20 units

```
. * Example from help file for user-written brain command
. clear
. set obs 200
number of observations (_N) was 0, now 200
. gen x = 4*_pi/200 *_n
. gen y = sin(x)
. brain define, input(x) output(y) hidden(20)
Defined matrices:
    input[4,1]
    output[4,1]
    neuron[1,22]
        1ayer[1,3]
        brain[1,61]
. quietly brain train, iter(500) eta(2)
. brain think ybrain
. sort x
. twoway (scatter y x) (lfit y x) (line ybrain x)
```


## Neural Networks Example (continued)

- We obtain



### 4.4 Further Details

- Chapter 10 of ISL Second Edition covers deep learning including
- convolutional neural networks for images
- document classification with e.g. word-pairs as features
- recurrent neural networks for e.g. autocorrelated time series
- detailed lab example using the keras package in Python
- https://www.statlearning.com/resources-second-edition has the same example done using the torch package in R .
- For deep learning a good text is
- Goodfellow, Yoshua Bengio and Aaron Courville (2016), Deep Learning, MIT Press.


## 5. Regression Trees and Random Forests: Overview

- Regression Trees sequentially split regressors $\mathbf{x}$ into regions that best predict $y$
- e.g., first split is income $<$ or $>\$ 12,000$ second split is on gender if income $>\$ 12,000$ third split is income $<$ or $>\$ 30,000$ (if female and income $>\$ 12,000$ ).
- Trees do not predict well
- due to high variance
- e.g. split data in two then can get quite different trees
- e.g. first split determines future splits (a greedy method).
- Better methods are then given
- bagging (bootstrap averaging) computes regression trees for different samples obtained by bootstrap and averages the predictions.
- random forests use only a subset of the predictors in each split
- boosting grows trees based on residuals from previous stage
- bagging and boosting are general methods (not just for trees).


### 5.1 Regression Trees

- Regression trees
- sequentially split $\mathbf{x}^{\prime} s$ into rectangular regions in way that reduces RSS
- then $\widehat{y}_{j}$ is the average of $y^{\prime} s$ in the region that $\mathbf{x}_{i}$ falls in
- with $J$ blocks RSS $=\sum_{j=1}^{J} \sum_{i \in R_{j}}\left(y_{i}-\bar{y}_{R_{j}}\right)^{2}$.
- Need to determine both the regressor $j$ to split and the split point $s$.
- For any regressor $j$ and split point $s$, define the pair of half-planes $R 1(j, s)=\left\{X \mid X_{j}<s\right\}$ and $R 2(j, s)=\left\{X \mid X_{j} \geq s\right\}$
- Find the value of $j$ and $s$ that minimize

$$
\sum_{i: \mathbf{x}_{i} \in R 1(j, s)}\left(y_{i}-\bar{y}_{R 1}\right)^{2}+\sum_{i: x_{i} \in R 2(j, s)}\left(y_{i}-\bar{y}_{R 2}\right)^{2}
$$

where $\bar{y}_{R 1}$ is the mean of $y$ in region $R 1$ (and similar for $R 2$ ).

- Once this first split is found, split both $R 1$ and $R 2$ and repeat
- Each split is the one that reduces RSS the most.
- Stop when e.g. less than five observations in each region.


## Tree example from ISL2 Figure 8.3 page 332

- (1) split X1 in two; (2) split the lowest X1 values on the basis of X2 into R1 and R2; (3) split the highest X1 values into two regions (R3 and R4/R5); (4) split the highest X1 values on the basis of X2 into R4 and R5.



## Tree example from ISL (continued)

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



## Regression tree (continued)

- The model is of form $f(\mathbf{x})=\sum_{j=1}^{J} c_{j} \times \mathbf{1}\left[\mathbf{x} \in R_{j}\right]$
- essentially OLS on a set of data-determined indicator variables.
- The approach is a topdown greedy approach
- top down as start with top of the tree
- greedy as at each step the best split is made at that particular step, rather than looking ahead and picking a split that will lead to a better tree in some future step
- so a seemingly worthless split early on in the tree might have been followed by a very good split later on.
- So deliberately overfit and then prune back
- use cost complexity pruning (or weakest link pruning)
- this adds a penalty in the number of terminal nodes and uses CV on this
- see ISL2 equation (8.4).


## Regression tree example

- This example is for duration data using Stata add-on cart
- I used it merely to illustrate what a tree looks like.



## Tree as alternative to k-NN or kernel regression

- Figure from Athey and Imbens (2019), "Machine Learning Methods Economists Should Know About"
- axes are $x_{1}$ and $x_{2}$
- note that tree used explanation of $y$ in determining neighbors
- tree may not do so well near boundaries of region
* random forests form many trees so not always at boundary.


Euclidean neighborhood, for KNN matching.


Tree-based neighborhood.

## 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 given next
- bagging
* bootstrap aggregating averages regression trees over many samples
- random forests
* additionally uses only a random subset of regressors at each split
- boosting
$\star$ trees build on preceding trees.


### 5.2 Bagging (Bootstrap Aggregating)

- Bagging is a general method for improving prediction that works especially well for regression trees.
- Idea is that averaging reduces variance (in nonlinear models).
- So average regression trees over many samples
- the different samples are obtained by bootstrap resample with replacement (so not completely independent of each other)
- for each sample obtain a large tree and prediction $\widehat{f}_{b}(\mathbf{x})$.
- average all these predictions: $\widehat{f}_{\text {bag }}(\mathbf{x})=\frac{1}{B} \sum_{b=1}^{B} \widehat{f}_{b}(\mathbf{x})$.
- Get test sample error by using out-of-bag (OOB) observations not in the bootstrap sample
- $\operatorname{Pr}\left[i^{\text {th }}\right.$ obs not in resample $]=\left(1-\frac{1}{n}\right)^{n} \rightarrow e^{-1}=0.368 \simeq 1 / 3$.
- this replaces cross validation.
- Interpretation of trees is now difficult so
- record the total amount that RSS is decreased due to splits over a given predictor, averaged over all $B$ trees.
- a large value indicates an important predictor.


### 5.3 Random Forests

- The $B$ bagging 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 use only a random sample of $m<p$ predictors in deciding each split (within each bootstrap sample)
- usually $m \simeq \sqrt{p}$
- this reduces correlation across bootstrap resamples
- simple bagging is random forest with $m=p$.
- Stata add-on command rforest implements random forests (and bagging)
- Matthias Schonlau and Rosie Zou (2020), "The random forest algorithm for statistical learning," The Stata Journal, 3-29.


## Random Forests (continued)

- Random forests are related to kernel and $k$-nearest neighbors
- as use a weighted average of nearby observations
- but with a data-driven way of determining which nearby observations get weight
- see Lin and Jeon (JASA, 2006).
- Susan Athey and coauthors are big on random forests.


### 5.4 Boosting

- Boosting is also a general method for improving prediction.
- Regression trees use a greedy algorithm.
- Boosting uses a slower algorithm to generate a sequence of trees
- each tree is grown using information from previously grown trees
- and is fit on a modified version of the original data set
- boosting does not involve bootstrap sampling.
- Specifically (with $\lambda$ a penalty parameter)
- given current model $b$ fit a decision tree to model $b^{\prime} s$ residuals (rather than the outcome $y$ )
- then update $\widehat{f}(\mathbf{x})=$ previous $\widehat{f}(\mathbf{x})+\lambda \widehat{f}^{b}(\mathbf{x})$
- then update the residuals $r_{i}=$ previous $r_{i}-\lambda \widehat{f}^{b}\left(\mathbf{x}_{i}\right)$
- the boosted model is $\widehat{f}(\mathbf{x})=\sum_{b=1}^{B} \lambda \widehat{f}^{b}\left(\mathbf{x}_{i}\right)$, a weighted sum of trees.
- Stata add-on boost includes file boost64.dll that needs to be manually copied into c: \ado $\backslash$ plus
- pylearn does trees, random forests and neural nets directly in Stata and requires installation of Python and the Python scikit-learn library.


### 5.5 Bayesian Additive Regression Trees

- Like boosting use only the original data
- whereas random forests draw random samples of the data.
- Start with $K$ trees and in each tree $\widehat{f}^{(1)}(\mathbf{x})=\bar{y}$.
- Then for each iteration $b=1, \ldots, B$
- for each tree $k$ and each observation $i$ create the partial residual which is $y_{i}$ minus the predictions from all other trees

$$
r_{i}=y_{i}-\sum_{k^{\prime}<k} \widehat{f}_{k}^{(b)}(\mathbf{x})-\sum_{k^{\prime}>k} \widehat{f}_{k}^{(b)}(\mathbf{x})
$$

- fit a new tree $\widehat{f}_{k}^{(b)}(\mathbf{x})$ to $r_{i}$ by randomly perturbing tree $k$ from the previous iteration $\widehat{f}_{k}^{(b-1)}(\mathbf{x})$ and favor perturbations that improve the fit
- compute $\widehat{f}_{k}^{(b)}(\mathbf{x})=\sum_{k=1}^{K} \widehat{f}_{k}^{(b-1)}(\mathbf{x})$
- Finally $\widehat{f}_{k}^{(b)}(x)=\frac{1}{B-L} \sum_{k=1}^{K} \widehat{f}_{k}^{(b-1)}(\mathbf{x})$ where $L$ is the number of burn-in reps.
- IT is an MCMC algorithm.
- ISL2 Section 8.2.4 has details.


## 6. Prediction Example

- Go through MUS2 Section 28.6 Prediction Example in detail
- Use code in ML_2022_part4.do with data set mus203mepsmedexp.dta
- Predict using 7 methods
- OLS with no interactions
- OLS with interactions
- LASSO with penalized coefficients
- Post LASSO (OLS with variables selected by LASSO)
- Neural network (add-on brain)
- Random forest (add-on randomforest)
- boost
- Fit on $80 \%$ of sample. See how predicts out of sample.


## Data

- Same MEPS data for 2013 on 65-90 year-olds. as in part 3.
- $y$ is ltotexp $=$ log total annual medical expenditure
- $\mathbf{x}$ is 5 continuous variables and 14 binary variables and $N=2955$
- same as part 3 except include suppins with other binary variables.
- With interactions get 188 unique variables.
. * Data for prediction example: 5 continuous and 14 binary variables
. qui use mus203mepsmedexp.dta, clear
. keep if ltotexp != .
(109 observations deleted)
. global xlist income educyr age famsze totchr
. global dlist suppins female white hisp marry northe mwest south ///
> msa phylim actlim injury priolist hvgg
. global rlist c.(\$xlist)\#\#c.(\$xlist) i.(\$dlist) c.(\$xlist)\#i.(\$dlist)
* Summary statistics for full sample . summarize ltotexp \$xlist \$dlist

| Variable | Obs | Mean | Std. Dev. | Min | Max |
| :---: | :---: | :---: | :---: | :---: | :---: |
| ltotexp | 2,955 | 8.059866 | 1.367592 | 1.098612 | 11.74094 |
| income | 2,955 | 22.68353 | 22.60988 | -1 | 312.46 |
| educyr | 2,955 | 11.82809 | 3.405095 | 0 | 17 |
| age | 2,955 | 74.24535 | 6.375975 | 65 | 90 |
| famsze | 2,955 | 1.890694 | . 9644483 | 1 | 13 |
| totchr | 2,955 | 1.808799 | 1.294613 | 0 | 7 |
| suppins | 2,955 | . 5915398 | . 4916322 | 0 | 1 |
| female | 2,955 | . 5840948 | . 4929608 | 0 | 1 |
| white | 2,955 | . 9736041 | . 1603368 | 0 | 1 |
| hisp | 2,955 | . 0812183 | . 2732163 | 0 | 1 |
| marry | 2,955 | . 5583756 | . 4966646 | 0 | 1 |
| northe | 2,955 | . 1536379 | . 3606623 | 0 | 1 |
| mwest | 2,955 | . 2318105 | . 42206 | 0 | 1 |
| south | 2,955 | . 3922166 | . 4883272 | 0 | 1 |
| msa | 2,955 | . 7397631 | . 438838 | 0 | 1 |
| phylim | 2,955 | . 4362098 | . 4959981 | 0 | 1 |
| actlim | 2,955 | . 2879865 | . 4529014 | 0 | 1 |
| injury | 2,955 | . 2020305 | . 4015828 | 0 | 1 |
| priolist | 2,955 | . 8240271 | . 3808616 | 0 | 1 |
| hvgg | 2,955 | . 6013536 | . 4897026 | 0 | 1 |

* OLS for full sample
. regress ltotexp \$xlist \$dlist, vce(robust) noheader

| ltotexp | Coef. | Robust <br> Std. Err. | t | $P>\|t\|$ | [95\% Conf. | Interval] |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| income | . 0007411 | . 0010967 | 0.68 | 0.499 | -. 0014092 | . 0028914 |
| educyr | . 0415116 | . 0076743 | 5.41 | 0.000 | . 0264641 | . 0565591 |
| age | . 0042834 | . 0037527 | 1.14 | 0.254 | -. 0030749 | . 0116416 |
| famsze | -. 0669498 | . 0261385 | -2.56 | 0.010 | -. 1182014 | -. 0156982 |
| totchr | . 3238205 | . 0188741 | 17.16 | 0.000 | . 2868126 | . 3608283 |
| suppins | . 1706101 | . 0469033 | 3.64 | 0.000 | . 0786434 | . 2625768 |
| female | -. 0508783 | . 0468787 | -1.09 | 0.278 | -. 1427968 | . 0410403 |
| white | . 1858472 | . 1325621 | 1.40 | 0.161 | -. 074077 | . 4457713 |
| hisp | -. 1101501 | . 0904202 | -1.22 | 0.223 | -. 2874435 | . 0671433 |
| marry | . 1751016 | . 0516199 | 3.39 | 0.001 | . 0738868 | . 2763164 |
| northe | . 2736686 | . 0713944 | 3.83 | 0.000 | . 1336804 | . 4136567 |
| mwest | . 3051208 | . 0689651 | 4.42 | 0.000 | . 169896 | . 4403456 |
| south | . 1957967 | . 0593267 | 3.30 | 0.001 | . 0794705 | . 3121229 |
| msa | . 0709307 | . 0512069 | 1.39 | 0.166 | -. 0294743 | . 1713357 |
| phylim | . 268737 | . 0567284 | 4.74 | 0.000 | . 1575054 | . 3799685 |
| actlim | . 3661458 | . 0636335 | 5.75 | 0.000 | . 241375 | . 4909165 |
| injury | . 1664688 | . 0539137 | 3.09 | 0.002 | . 0607564 | . 2721813 |
| priolist | . 4361775 | . 0689187 | 6.33 | 0.000 | . 3010436 | . 5713114 |
| hvgg | -. 0959803 | . 0463345 | -2.07 | 0.038 | -. 1868316 | -. 0051289 |
| _cons | 5.633868 | . 3425158 | 16.45 | 0.000 | 4.962272 | 6.305463 |

- We will use $80 \%$ of sample for training and remaining $20 \%$ for out-of-sample evaluation
. * Split the sample with $80 \%$ in training sample
. splitsample ltotexp, generate(train) split(1 4) values(0 1) rseed(10101)
. tabulate train

| train | Freq. | Percent | Cum. |
| ---: | ---: | ---: | ---: |
| 0 | 591 | 20.00 | 20.00 |
| 1 | 2,364 | 80.00 | 100.00 |
| Total | 2,955 | 100.00 |  |

- Then go through code in ML_2022_part4.do


## Predictions from Various Models

* OLS with 19 regressors
regress Itotexp \$xlist \$dlist if train==1, noheader vce(robust)
qui predict y_small
* OLS with 188 potential regressors and 104 estimated
qui regress Itotexp \$rlist if train==1
qui predict y_full
* LASSO with 188 potential regressors leads to 32 selected
qui lasso linear Itotexp \$rlist if train==1, selection(adaptive) rseed(10101) nolog
qui predict y_laspen // use penalized coefficients
qui predict y_laspost, postselection // use post selection OLS coeffs


## Predictions from Various Models (continued)

* Principal components using the first 5 principal components of 19 variables qui pca \$xlist \$dlist if train==1
qui predict pc *
qui regress Itotexp pc1-pc5 if train==1
qui predict y_pca
* Neural network: 19 variables one hidden layers with 10 units
* This did not work on my latest computer
brain define, input(\$xlist \$dlist) output(Itotexp) hidden(10)
qui brain train if train $==1$, iter(500) eta(2) // eta $>1$ uses SGD
brain think y_neural
* Random forest with 19 variables
qui rforest Itotexp \$xlist \$dlist if train==1, ///
type(reg) iter(200) depth(10) Isize(5)
qui predict y_ranfor


## Predictions from Various Models (continued)

- Compute training MSE and test MSE

```
foreach var of varlist y_noreg y_small y_full y_laspen ///
    y_laspost y_pca y_neural y_ranfor {
    qui gen 'var'errorsq = ('var' - Itotexp)^2
    qui sum 'var'errorsq if train == 1
    scalar mse'var'train =r(mean)
    qui sum 'var'errorsq if train == 0
    qui scalar mse'var'test = r(mean)
    display "Predictor:" "'var'" _col(21) ///
        " Train MSE = " %5.3f mse'var'train ///
        " Test MSE = " %5.3f mse'var'test
    }
```


## Predictions from Various Models (continued)

- Training sample: Flexible models - random forest and neural networks did best.
- Test sample: Simpler models - LASSO and small model OLS did best.

| Predictor: $y$ y_noreg | Train MSE $=1.821$ | Test MSE $=2.063$ |
| :--- | :--- | :--- |
| Predictor: $y$ ysmall | Train MSE $=1.339$ | Test MSE $=1.492$ |
| Predictor: y_full | Train MSE $=1.262$ | Test MSE $=1.509$ |
| Predictor: $y$ y_laspen | Train MSE $=1.298$ | Test MSE $=1.491$ |
| Predictor: y_laspost | Train MSE $=1.297$ | Test MSE $=1.493$ |
| Predictor: y_pca | Train MSE $=1.397$ | Test MSE $=1.545$ |
| Predictor: y_neural | Train MSE $=1.211$ | Test MSE $=1.808$ |
| Predictor: y_ranfor | Train MSE $=1.047$ | Test MSE $=1.574$ |

## 7. Prediction for Economics

- Hal Varian (2014) has early survey.
- Mullainathan and Spiess (2017)
- summarizes various
- has good application to housing prices(already presented)
- has good summary of recent economics ML applications.


### 7.1 Hal Varian 2014 Survey

- Hal Varian (2014), "Big Data: New Tricks for Econometrics," JEP, Spring, 3-28.
- Surveys tools for handling big data
- file system for files split into large blocks across computers
* Google file system (Google), Hadoop file system
- database management system to handle large amounts of data across many computers
* Bigtable (Google), Cassandra
- accessing and manipulating big data sets across many computers
^ MapReduce (Google), Hadoop.
- language for MapReduce / Hadoop
^ Sawzall (Google), Pig
- Computer language for parallel processing
* Go (Google - open source)
- simplified structured query language (SQL) for data enquiries
^ Dremel, Big Query (Google), Hive, Drill, Impala.


## Hal Varian (continued)

- Surveys methods
- article discusses k-fold CV, trees, lasso, ....
- small discussion of causality and prediction
- (note that a classic fail is Google flu trends)
- for references mentions ESL and ISL.


### 7.2 Summary of Machine Learning Algorithms

- From Mullainathan and Spiess (2017)


## Table 2

## Some Machine Learning Algorithms

| Function class $\mathcal{F}$ ( and its parametrization) | Regularizer $R(f)$ |
| :---: | :---: |
| Global/parametric predictors <br> Linear $\beta^{\prime} x$ (and generalizations) | Subset selection $\\|\beta\\|_{0}=\sum_{j=1}^{b} 1_{\beta_{j} \neq 0}$ <br> LASSO $\\|\beta\\|_{1}=\sum_{j=1}^{k}\left\|\beta_{j}\right\|$ <br> Ridge $\\|\beta\\|_{2}{ }^{2}=\sum_{j=1}^{k} \beta_{j}{ }^{2}$ <br> Elastic net $\alpha\\|\beta\\|_{1}+(1-\alpha)\\|\beta\\|_{2}{ }^{2}$ |
| Local/nonparametric predictors Decision/regression trees | Depth, number of nodes/leaves, minimal leaf size, information gain at splits |
| Random forest (linear combination of trees) | Number of trees, number of variables used in each tree, size of bootstrap sample, complexity of trees (see above) |
| Nearest neighbors | Number of neighbors |
| Kernel regression | Kernel bandwidth |

## Table 2 (continued)

## Mixed predictors

Deep learning, neural nets, convolutional neural networks

Splines

## Combined predictors

Bagging: unweighted average of predictors from bootstrap draws

Boosting: linear combination of predictions of residual

Ensemble: weighted combination of different predictors

Number of levels, number of neurons per level, connectivity between neurons

Number of knots, order

Number of draws, size of bootstrap samples (and individual regularization parameters)

Learning rate, number of iterations (and individual regularization parameters)
Ensemble weights (and individual regularization parameters)

### 7.3 Some Thoughts on ML Prediction

- Clearly there are many decisions to make in implementation
- how are features converted into x's
- tuning parameter values
- which ML method to use
- even more with an ensemble forecast.
- For commercial use this may not matter
- all that matters is that predict well enough.
- But for published research we want reproducibility
- At the very least document exactly what you did
- provide all code (and data if it is publicly available)
- keep this in mind at the time you are doing the project.
- For public policy we prefer some understanding of the black box
- this may be impossible.


## 8. Software for Machine Learning

- This list will change over time and is not necessarily the best
- There are over 29,000 R packages in CRAN and over 137,000 Python packages.
- Python libraries include
- scikit-learn for most ML methods including trees and random forests and basic neural networks
- keras and tensorflow for neural networks.
- R packages used in ISLR2 include
- spline library for splines
- torch package for neural networks
- tree library and randomForest package for tree-based methods.
- Stata commands include
- npregress for local regression, series regression and splines
- pca for principal components


## Software (continued)

- Stata add-ons include
- brain for neural networks (very basic)
- rforest for trees and random forests
- crtrees for trees and random forests
- srtrees wrapper for R commands for trees and random forests
- r_ml_stata.ado and r_ml_stata.ado wrappers for ML methods including neural networks and trees in the python scikit-learn library

ฝ https://sites.google.com/view/giovannicerulli/machine-learning-instata

- pylearn is a wrapper for trees and random forests in python scikit-learn library


## 9. References

- ISL2: Gareth James, Daniela Witten, Trevor Hastie and Robert Tibsharani (2021), Second Edition, An Introduction to Statistical Learning: with Applications in $R$, Springer.
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- second edition coming soon.
- ESL: Trevor Hastie, Robert Tibsharani and Jerome Friedman (2009), The Elements of Statistical Learning: Data Mining, Inference and Prediction, Springer.
- free legal pdf at http://statweb.stanford.edu/~tibs/ElemStatLearn/index.html
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- EH: Bradley Efron and Trevor Hastie (2016), Computer Age Statistical Inference: Algorithms, Evidence and Data Science, Cambridge University Press.
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