Supervised Learning (Part I)

DS 6030 | Fall 2022

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1 Supervised Learning Intro

1.1 Survey

There will be a survey at end of lecture. You can access the survey from MS Teams. At the top tab of the class find "Fill | Supervised Learning I: What is the best model?". You may need to expand "More".

1.2 Required R Packages

We will be using the R packages of:

- FNN for k nearest neighbor models
- tidyverse for data manipulation and visualization
- broom for tidying model output

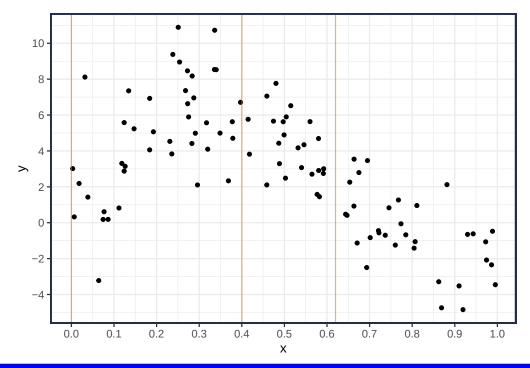
```
library(FNN)
library(broom)
library(tidyverse)
```

1.3 Supervised Learning

- In *supervised learning*, each observation can be partitioned into two sets: the predictor variables and the outcome variable(s).
 - Predictor variables are sometimes called independent/feature variables
 - Outcome variables are sometimes called target/labels/response/dependent variables.
- \bullet Usually the predictor variables are represented by X and the response variables represented by Y
- The goal in supervised learning is to find the patterns and relationships between the predictors, X, and the response, Y.
 - Usually the goal is to *predict* the value of Y given X.
- Later in the course we will explore the *unsupervised learning* topics of association analysis, network analysis, density estimation, clustering, and anomaly detection which do not have any outcomes (i.e., no Y's).

2 Example Data

Consider some data $D = \{(X_i, Y_i)\}_{i=1}^n$ with $Y_i \in \mathbb{R}$, $X_i \in [0, 1]$ and n = 100.



Your Turn #1

The goal is to predict new Y values if we are given the X's.

- If x = .40, predict Y.
- If x = 0, predict Y.
- If x = .62, predict Y.
- How should we build a *model* that will automatically predict Y for any given X?

3 Linear Models

• <u>Linear models</u> refer to a class of models where the output (predicted value) is a linear combination (weighted sum) of the input variables

$$f(x;\beta) = \beta_0 + \sum_{j=1}^{p} \beta_j x_j$$

where $x = [x_1, \dots, x_p]^\mathsf{T}$ is a vector of features/variables/attributes and $\hat{Y}|x = f(x; \hat{\beta})$ is the predicted response at X = x

- the coefficients (or weights), $\hat{\beta}$ are often selected by minimizing the squared residuals of the *training* data (may also be described as *ordinary least squares*)
 - But, there are other, and better, ways to estimate the parameters in linear regression that we will discuss later in the course. (e.g., Lasso, Ridge, Robust)

3.1 Simple Linear Regression

• single predictor variable $x \in \mathbb{R}$

- $f(x;\beta) = \beta_0 + \beta_1 x$
- Use training data: $D_{\text{train}} = \{(x_i, y_i)\}_{i=1}^n$
- OLS uses the weights/coefficients that minimize the RSS loss function over the training data

$$\hat{\beta} = \underset{\beta}{\operatorname{arg\,min}} \operatorname{SSE}(\beta)$$

• where SSE is the sum of squared errors (also known as residual sum of squares (RSS))

$$SSE(\beta) = \sum_{i}^{n} (y_i - f(x_i, \beta))^2$$

$$= \sum_{i}^{n} (y_i - \beta_0 - \beta_1 x_i)^2$$

$$= \sum_{i}^{n} \hat{\epsilon}_i^2 \quad \text{where } \hat{\epsilon}_i = y_i - \hat{y}_i \text{ is the residual}$$

· The solutions are

$$\hat{\beta}_0 = \bar{y} - \beta_1 \bar{x}$$

$$\hat{\beta}_1 = \frac{\sum_{i=1}^n (x_i - \bar{x})(y_i - \bar{y})}{\sum_{i=1}^n (x_i - \bar{x})^2}$$

• Definitions:

$$MSE(\beta) = \frac{1}{n}SSE(\beta)$$

$$= \frac{1}{n} \sum_{i=1}^{n} (y_i - f(x_i; \beta))^2$$

$$RMSE = \sqrt{MSE} = \sqrt{SSE} / \sqrt{n}$$

3.2 OLS Linear Models in R

3.2.1 Estimation with 1m()

In **R**, the function lm() fits an OLS linear model

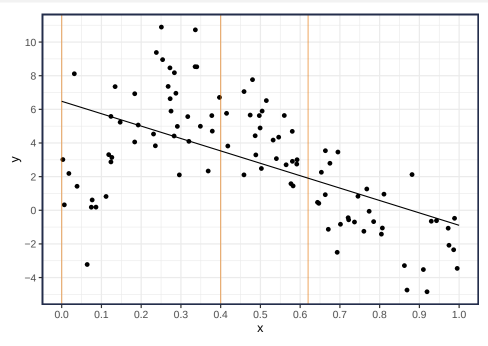
```
#> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
#>
#> Residual standard error: 2.91 on 98 degrees of freedom
#> Multiple R-squared: 0.331, Adjusted R-squared: 0.325
#> F-statistic: 48.6 on 1 and 98 DF, p-value: 3.69e-10
broom::tidy(m1) # model coefficients (as a data frame)
#> # A tibble: 2 x 5
#> term estimate std.error statistic p.value
#> <chr> <dbl> <dbl> <dbl> <dbl> <dbl> = 1 (Intercept) 6.48 0.584 11.1 5.39e-19
                -7.37 1.06 -6.97 3.69e-10
#> 2 x
broom::glance(m1)
                       # model properties
#> # A tibble: 1 x 12
#> # ... with 2 more variables: df.residual <int>, nobs <int>, and abbreviated
#> # variable names 1: adj.r.squared, 2: statistic, 3: deviance
#> # i Use `colnames()` to see all variable names
```

- lm() uses the formula interface, which includes the intercept by default.
 - Some examples of using formulas as well as getting the underlying X (model/design matrix) can be found here

3.2.2 Prediction with predict ()

The function predict () is used to get the predicted values.

```
xseq = seq(0, 1, length=200)  # sequence of equally spaced values from 0 to 1
xeval = tibble(x = xseq)  # make into a tibble object
yhat1 = predict(m1, xeval)  # vector of yhat's (predictions)
```



3.2.3 Questions

Your Turn #2

- 1. How did we do? If $X_{\rm new}$ is close to 0, or close to 0.4, or close to .62?
- 2. How to make it better?

4 Polynomial inputs

- In the *simple* linear regression model, we had 2 parameters that we needed to estimation, β_0 and β_1 . Thus, the model complexity is minimal.
 - The only thing simpler is an intercept only model.
- But the data appears to have a more *complex* structure than linear.
- A parametric approach to add complexity is to incorporate polynomial terms into the model.
 - A quadratic model is $f(x; \beta) = \beta_0 + \beta_1 x + \beta_2 x^2$

4.1 Estimation

• OLS uses the weights/coefficients that minimize the SSE loss function over the training data

$$\hat{\beta} = \underset{\beta}{\operatorname{arg\,min}} \quad \operatorname{SSE}(\beta) \qquad \text{Note: } \beta \text{ in this problem is a } \textit{vector}$$

$$= \underset{\beta}{\operatorname{arg\,min}} \quad \sum_{i=1}^{n} (y_i - f(x_i; \beta))^2$$

$$= \underset{\beta}{\operatorname{arg\,min}} \quad \sum_{i=1}^{n} (y_i - \beta_0 - \beta_1 x_i - \beta_2 x_i^2)^2$$

4.1.1 Matrix notation

Model

$$f(\mathbf{x}; \beta) = \mathbf{x}^\mathsf{T} \beta$$

$$\mathbf{x} = \begin{bmatrix} 1 \\ x \\ x^2 \end{bmatrix} \qquad \beta = \begin{bmatrix} \beta_0 \\ \beta_1 \\ \beta_2 \end{bmatrix}$$

Your Turn #3: Matrix Notation

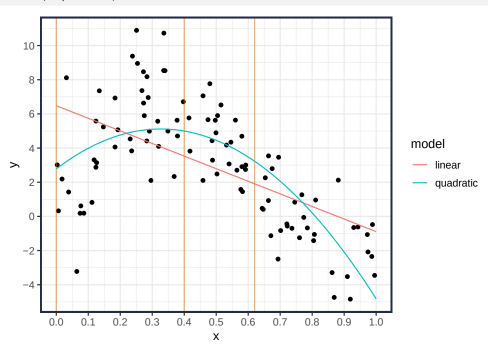
Solve for $\hat{\beta}$ using matrix notation.

$$Y = \begin{bmatrix} Y_1 \\ Y_2 \\ \vdots \\ Y_n \end{bmatrix} \qquad X = \begin{bmatrix} 1 & X_1 & X_1^2 \\ 1 & X_2 & X_2^2 \\ \vdots & \vdots & \vdots \\ 1 & X_n & X_n^2 \end{bmatrix} \qquad \beta = \begin{bmatrix} \beta_0 \\ \beta_1 \\ \beta_2 \end{bmatrix}$$

4.1.2 R implementation

In \mathbf{R} , the function poly () is a convenient way to get polynomial terms

```
m2 = lm(y~poly(x, degree=2), data=data_train)
yhat2 = predict(m2, xeval)
```



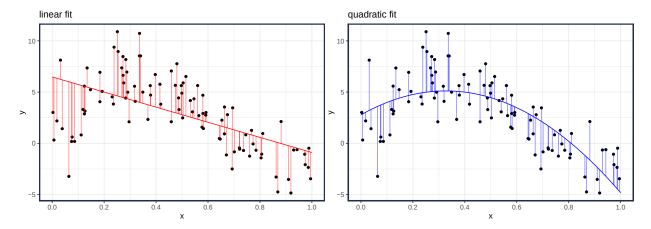
Your Turn #4

- 1. How did we do? If $X_{\rm new}$ is close to 0, or close to 0.4, or close to .62?
- 2. But does the quadratic model fit better *overall*?
- 3. What is the *complexity* of the quadratic model?

4.2 Performance Comparison (on Training Data)

Comparing the two models (according to MSE), the quadratic model does much better!

degree	MSE	npars
1	8.29	2
2	5.58	3



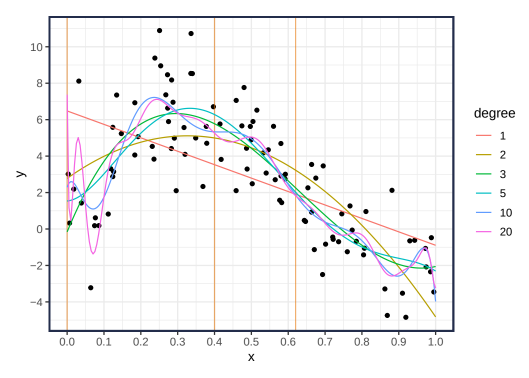
As my kids always reason, "if a little is good, than a lot must be better". So why not try more complex models by increasing the polynomial degree.

ullet Polynomial of degree d

$$f_{\text{poly}}(x; \beta, d) = \beta_0 + \sum_{j=1}^{d} \beta_j x^j$$

degree	MSE	npars
1	8.29	2
2	5.58	3
3	4.28	4
5	4.10	6
10	3.65	11
20	3.16	21

And its always good to observe the plot



- For degree=20, the behavior at the end points are a bit erratic.
- Using a higher degree would further reduce the MSE, but the fitted curve would be more "complex" and may not be as good for new data.

5 k-nearest neighbor models

- The k-NN method is a non-parametric *local* method, meaning that to make a prediction $\hat{y}|x$, it only uses the training data in the *vicinity* of x.
 - contrast with OLS linear regression, which uses all X's to get prediction.
- The model is simple to describe

$$f_{knn}(x;k) = \frac{1}{k} \sum_{i:x_i \in N_k(x)} y_i$$
$$= \text{Avg}(y_i \mid x_i \in N_k(x))$$

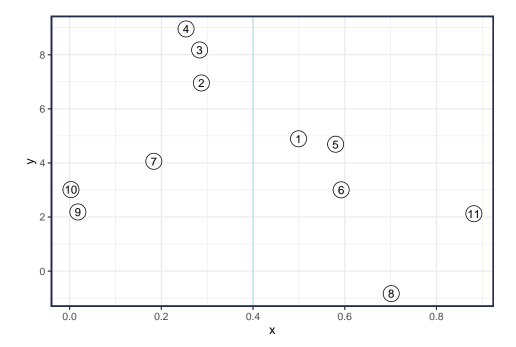
- $N_k(x)$ are the set of k nearest neighbors to x
- only the k closest y's are used to generate a prediction
- it is a *simple mean* of the k nearest observations

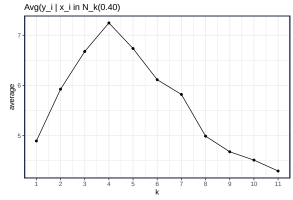
Your Turn #5

What is the estimate $f_{knn}(x; k = n)$?

5.0.1 Example

Consider the following example where we wish to estimate $Y \mid X = 0.40$





X	у	k	D	$\hat{f}_{\mathrm{knn}}(x;k)$
0.50	4.89	1	0.10	4.89
0.29	6.96	2	0.11	5.92
0.28	8.18	3	0.12	6.68
0.25	8.95	4	0.15	7.25
0.58	4.69	5	0.18	6.73
0.59	3.00	6	0.19	6.11
0.18	4.06	7	0.22	5.82
0.70	-0.83	8	0.30	4.99
0.02	2.19	9	0.38	4.68
0.00	3.01	10	0.40	4.51
0.88	2.12	11	0.48	4.29

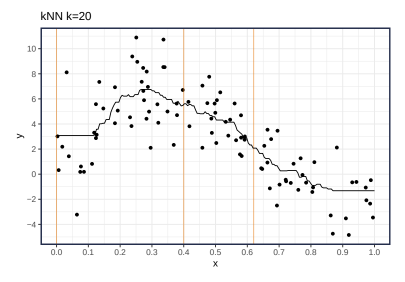
5.0.2 Notes about knn

- A suitable *distance* measure (e.g. Euclidean) must be chosen.
 - And predictors are often scaled (same sd or range) so one variable doesn't dominate the distance calculation
- Because the distance to neighbors grows exponentially with increased dimensionality/features, the *curse of dimensionality* is often referenced with respect to knn.
 - This means that in high dimensions most *neighbors* are not very close and the method becomes less *local*
- One computational drawback of knn methods is that all the training data must be stored in order to make predictions.
 - For large training data, may need to sample (or use prototypes)

5.1 knn in action

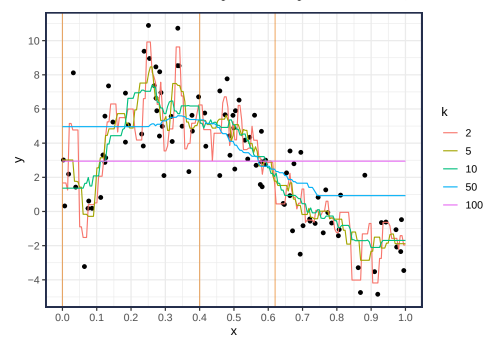
In ${\bf R}$, the function knn.reg() from the FNN package will fit a knn regression model. Here is a k=20 nearest neighbor model

```
library(FNN) # library() loads the package. Access to knn.reg()
#- fit a k=20 knn regression
knn.20 = knn.reg(select(data_train, x), test=xeval, y=data_train$y, k=20)
```



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- The complexity of a knn model increases as k decreases.
- The least complex model, which is a constant, occurs when k=n
- The most complex model when k=1
- The effective degrees of freedom or edf for a knn model is n/k
 - this is a measure of the model *complexity*. It is approximately the number of parameters that are estimated in the model (to allow comparison with parametric models)



5.1.1 Performance of the knn models (on training data)

k	MSE	edf
100	12.40	1
50	6.87	2
10	3.86	10
5	3.16	20
2	1.84	50

6 Predictive Model Comparison (or how to choose the best model)

6.1 Predictive Model Evaluation

Our goal is prediction, so we should evaluate the models on their *predictive performance*.

- We need to use hold-out data (i.e., data not used to fit the model) to evaluate how well our models do in prediction
- Call these data test data $D_{\text{test}} = \{(X_j, Y_j)\}_{j=1}^J$
 - Note: assume that the test data comes from the same distribution as the training data
 - Or $P_{\text{test}}(X, Y) = P_{\text{train}}(X, Y)$
 - both Y and X from same distribution
- Later in the course we will cover ways to do this when we only have training data (e.g., cross-validation)
- but for today, we have an unlimited amount of *test data* at our disposal (since we know how the data were generated)

6.2 Statistical Decision Theory

- In a prediction context, we want a *point estimate* for the value of an unobserved r.v. $Y \in \mathbb{R}$ given an input feature $X \in \mathbb{R}$.
- Let f(X) be the prediction of Y given X.
- Define a loss function L(Y, f(X)) that indicates how bad it is if we estimate the value Y by f(X)
 - E.g. Y is the number of customers complaints in a call center and X is the day of week

 $L(Y, f(X)) = (Y - f(X))^2$

- If we guess f(X) = 500, but there are really Y = 2000, how bad would that be?
- A common loss function is squared error

Y - f(X)

• The best model is the one that minimizes the *expected loss* or Risk or Expected Prediction Error (EPE)

$$Risk = EPE = E[loss]$$

• For *squared error*, the *risk* for using the model *f* is:

$$R(f) = E_{XY}[L(Y, f(X))]$$

= $E_{XY}[(Y - f(X))^{2}]$

where the expectation is w.r.t. the *test values* of X, Y.

- Note under squared error loss, the risk is also known as the mean squared error (MSE)
- To simplify a bit, let's examine the risk of model f at a given fixed input X = x. This removes the uncertainty in X, so we only have uncertainty coming from Y.

$$R_x(f) = E[L(Y, f(x)) \mid X = x]$$

= $E[(Y - f(x))^2 \mid X = x]$ for squared error loss

where the expectation is taken with respect to Y|X=x

• The best prediction $f^*(x)$, given X = x, is the value that minimizes the risk

$$f^*(x) = \underset{c}{\arg\min} R_x(c)$$
$$= \underset{c}{\arg\min} \operatorname{E}[(Y - c)^2 \mid X = x]$$

Your Turn #6

What is the optimal prediction at X = x under the squared error loss?

• I.e., find $f^*(x)$.

6.2.1 Squared Error Loss Functions

- Conclusion: If quality of prediction is measured by squared error, then the best predictor is the (conditional) expected value $f^*(x) = E[Y|X=x]$.
 - And the minimum Risk/MSE is $R_x(f^*) = V[Y|X=x]$
- Summary: Under squared error loss the Risk (at input x) is

$$R_x(f) = \mathrm{E}_Y[L(Y,f(X)) \mid X = x]$$

$$= \mathrm{E}_Y[(Y - f(x))^2 \mid X = x] \qquad \text{using squared error loss}$$

$$= \mathrm{V}[Y \mid X = x] + (E_Y[Y \mid X = x] - f(x))^2$$

$$= \mathrm{Irreducible \ Variance} + \mathrm{model \ squared \ error}$$

6.2.2 kNN and Polynomial Regression

• The kNN model estimates the conditional expectation by using the data in a *local region* around x

$$\hat{f}_{knn}(x;k) = Ave(y_i \mid x_i \in N_k(x))$$

This assumes that the true f(x) can be well approximated by a *locally constant* function

• Polynomial (linear) regression, on the other hand, assumes that the true f(x) is well approximated by a globally polynomial function

$$\hat{f}_{\text{poly}}(x;d) = \beta_0 + \sum_{j=1}^{d} \beta_j x^j$$

6.2.3 Empirical Risk

• The actual Risk/EPE is based on the expected error from *test data* (out-of-sample), or data that was not used to estimate \hat{f}

$$R(f) = E_{XY}[L(Y, f(X))]$$

$$= E_{XY}[(Y - f(X))^{2}]$$
 for squared error loss

where X, Y are from Pr(X, Y) (i.e., test data)

• But is it a bad idea to choose the best model according to empirical risk or training error?

$$\hat{R}_n(f) = \frac{1}{n} \sum_{i=1}^n L(y_i, f(x_i))$$

$$= \frac{1}{n} \sum_{i=1}^n (y_i - f(x_i))^2$$
 for squared error loss

6.3 Choose the best *predictive* model

Your Turn #7

Which model will you choose?

Enter your answer on MS Teams. At the top tab of the class find "Fill | Supervised Learning I : What is the best model?". You may need to expand "More".

