PDAT615G: Machine Learning

Module 1 - Introduction

/ DISTINCT BY DESIGN

Module 1: Introduction

- Supervised, unsupervised and reinforcement learning
- Regression models
 - Single/multiple variable for numeric prediction
 - Logistic regression for classification
- Review of R programming techniques
 - Defining functions
 - Vectorized commands & loop-like constructions

Types of Machine Learning

Supervised learning trains a model to use certain features, or variables, to predict a response.

- Training data contains both explanatory variables and the response.
- Models are trained to be able to use only the explanatory variables to predict new responses.
- Regression is an example of a supervised learning technique.

Unsupervised learning analyzes patterns in data without a particular response variable specified.

- Clustering is one example. What groupings appear in the data?
- Dimension reduction is another. Do fewer variables describe the data just as well?

Reinforcement learning creates an algorithm by rewarding "good" behavior and punishing "bad" behavior.

- Reinforcement learners need
 - a way to measure the desirability of the end state,
 - methods to explore the space of all possible actions, and
 - · methods to find optimal strategies.
- Examples might include
 - a game-playing algorithm, or
 - a planner for supply-chain management.

Regression Models

Linear regression models can be used to predict a numeric response variable.

Linear regression models predict the value of one variable (the response, or dependent variable) based on other variables (the predictor, or independent, variables):

$$Y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \cdots + \beta_k x_k + \varepsilon.$$

Response (dependent) variable: denoted Y

The response variable Y must be a numeric variable.

Explanatory (independent) variables: denoted $x_1, x_2, ..., x_k$ Predictors $x_1, ..., x_k$ can be numeric or categorical variables.

If we have **MORE THAN ONE** independent variable, say x_1 and x_2 , or x_1 , x_2 and x_3 , the model is a *multiple linear regression model*.

Regression coefficients are chosen to minimize the sum of squared residuals.

■ We assume a model form:

$$Y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \cdots + \beta_k x_k + \varepsilon.$$

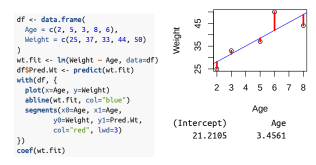
■ The β 's are estimated by b_0, b_1, \ldots , giving the regression equation to calculate \hat{y} , the predicted value of y.

$$\hat{y} = b_0 + b_1 x_1 + b_2 x_2 + \dots + b_k x_k.$$

■ The coefficients b_0, b_1, \ldots, b_k are chosen to minimize

$$\sum_{i=1}^n (y_i - \hat{y}_i)^2$$

Regression models are calculated with lm, and predicted values obtained with predict.



Logistic regression predicts the likelihood of a Yes/No categorical variable.

- Suppose *p* represents the probability of a "Yes" response.
- lacksquare Perhaps p is related to one or more predictor variables.
- Logistic regression models the log odds as a linear function of one or more variables:

$$\log\left(\frac{\rho}{1-\rho}\right) = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \dots + \varepsilon, \text{ or } \rho = \frac{1}{1+e^{-(\beta_0+\beta_1 x_1 + \dots + \varepsilon)}}$$

■ Note: Functions of the form $p=\frac{1}{1+e^{-z}}$ asymptotically approach 0 and 1 in an "S"-shaped curve, which is good for modeling probabilities.

Logistic regression models are calculated with the glm command and family="binomial".

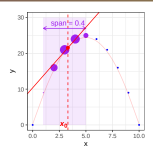
```
# Note: glm accepts 0/1 or
                                                0.8
# factor variables.
df$Vax <- c(0, 1, 0, 1, 0)
                                                0.4
vax.fit \leftarrow glm(Vax \sim Age, data=df,
                family="binomial")
with(df, plot(x=Age, y=Vax))
# Plot the curve
                                                                 5
                                                                     6 7 8
df.curve <- data.frame(
  Age = seq(2, 8, by=0.01)
                                                                Age
df.curve$p <- predict(vax.fit,</pre>
  newdata=df.curve, type="response")
with(df.curve,
 lines(x=Age, y=p, col="blue"))
```

LOESS creates a "local" regression model near each point by using only near points in model construction.

- Historically, two related terms have been used.
- LOWESS LOcally Weighted Scatterplot Smoothing
 - lowess(): Calculates a fit for one *x* and one *y*. Outputs predicted points for plotting.
 - · Defaults to first-degree (linear) fitting.
- LOESS LOcally Estimated Scatterplot Smoothing
 - Interface similar to lm().
 - Outputs a model for use with predict().
 - Generalizes LOWESS. Can accommodate multiple predictor variables.
 - · Defaults to second-degree polynomial fitting.

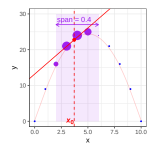
Example: Degree 1 LOESS Fit, 40% Span

- We want to find a function $\hat{f}(x)$ that fits the data.
- Step 1: For a base point x_0 , find the closest 40% of the data points.
- Step 2: Weight each point within the span according to its distance from x₀.
- Step 3: Calculate a weighted linear regression using those points.
- Step 4: $\hat{f}(x_0)$ is the point on that line above x_0
- Step 5: Repeat for other x's.



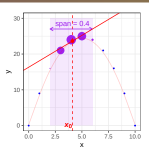
Example: Degree 1 LOESS Fit, 40% Span

- We want to find a function $\hat{f}(x)$ that fits the data.
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- Step 5: Repeat for other x's.



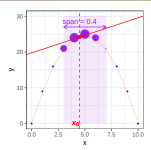
Example: Degree 1 LOESS Fit, 40% Span

- We want to find a function $\hat{f}(x)$ that fits the data.
- Step 1: For a base point x₀, find the closest 40% of the data points.
- Step 2: Weight each point within the span according to its distance from x₀.
- Step 3: Calculate a weighted linear regression using those points.
- Step 4: $\hat{f}(x_0)$ is the point on that line above x_0
- Step 5: Repeat for other x's.



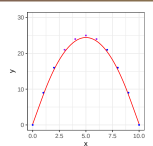
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- Step 4: $\hat{f}(x_0)$ is the point on that line above x_0
- Step 5: Repeat for other x's.

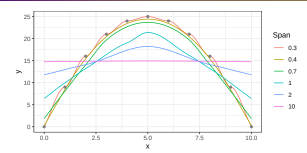


Example: Degree 1 LOESS Fit, 40% Span

- We want to find a function f(x) that fits the data.
- Step 1: For a base point x_0 , find the closest 40% of the data points.
- Step 2: Weight each point within the span according to its distance from x₀.
- Step 3: Calculate a weighted linear regression using those points.
- Step 4: $\hat{f}(x_0)$ is the point on that line above x_0
- Step 5: Repeat for other x's.



Increasing the span moves from a more "local" to a more "global" fit.



 $\begin{tabular}{l} \textbf{Note:} Increasing span larger than 1 considers all the data points and evens out the weights, approaching a standard linear regression. \end{tabular}$

Evaluating Predictive Models

Mean squared error (MSE) or root mean squared error (RMSE) are often used with numeric predictors.

MSE =
$$\frac{1}{n} \sum_{i=1}^{n} (y - \hat{y})^2$$

$$\mathsf{RMSE} = \sqrt{\frac{1}{n} \sum_{i=1}^n (y - \hat{y})^2}$$

RMSE is measured in the same units as the data.

```
df

Age Weight Pred.Wt Vax

1 2 25 28.123 0

2 5 37 38.491 1

3 3 33 31.579 0

4 8 44 48.860 1

5 6 50 41.947 0

sqrt(mean((df$Weight - df$Pred.Wt)^2))

[1] 4.5267

RMSE(pred=df$Pred.Wt, obs=df$Weight)

[1] 4.5267
```

Accuracy, Sensitivity and Specificity give information about classifiers.

	Act. Pos	Act. Neg
Pr. Pos	Α	В
Pr. Neg	С	D

Accuracy Percentage of correct predictions. $\left(\frac{A+D}{A+B+C+D}\right)$

Sensitivity Percentage of actual positives correctly predicted to be positive. $\left(\frac{A}{A+C}\right)$

Specificity Percentage of actual negatives correctly predicted to be negative. $\left(\frac{D}{B+D}\right)$

```
df$Prob.Vax <-
    predict(vax.fit, type="response")
df$Pred.Vax <-
    as.numeric(df$Prob.Vax > 0.50)
# Accuracy
mean(df$Vax == df$Pred.Vax)
[1] 0.6
# Sensitivity
with(df[df$Vax==1, ],
    mean(Vax == Pred.Vax))
[1] 0.5
# Specificity
with(df[df$Vax==0, ],
    mean(Vax == Pred.Vax))
[1] 0.66667
```

For classifiers that use a probability cut-off, the ROC curve describes the trade-off between sensitivity and specificity.

- For a set of thresholds between 0 and 1, predict the binary response.
- Calculate the pair (Specificity, Sensitivity).
- Plot the point on the graph.
- When AUC ≈ 1, a threshold can be chosen to make both sensitivity and specificity high.

Vax Prob.Vax 1 0 0.038597 2 0 0.090001 3 1 0.375094 4 0 0.596565 5 1 0.899743

```
library(pROC)
vax.roc <- roc(Vax ~ Prob.Vax, data=df)
plot(vax.roc, type="b")

2.0.8 0.4 0.0
Specificity

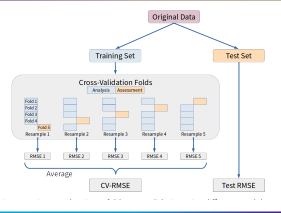
auc(vax.roc)
```

Area under the curve: 0.833

Predictive models should be evaluated on data not used to train the model.

```
Simplest: Create training and
                                       # Caret's `createDataPartition`
testing sets.
                                       # creates indices as well.
# Base R: The `sample` command
                                       # Given a factor, it will try to
# selects random indices.
                                       # split them evenly.
ind <- sample(1:nrow(df),</pre>
                                       ind <- createDataPartition(df$Vax,</pre>
        round(0.70*nrow(df)))
                                                        p=0.7, list=FALSE)
df.train <- df[ind, ]
df.test <- df[-ind, ]</pre>
                                       df.train <- df[ind, ]
                                       df.test <- df[-ind, ]</pre>
df.train[, 1:4]
                                       \texttt{df.train[, 1:4]}
  Age Weight Pred.Wt Vax
                                         Age Weight Pred.Wt Vax
          33 31.579 0
                                                 25 28.123
          50 41.947
                                                 37 38.491
          25 28.123
                                                 44 48.860
          44 48.860
                                                 50 41.947
df.test[, 1:4]
                                       df.test[, 1:4]
  Age Weight Pred.Wt Vax
                                         Age Weight Pred.Wt Vax
2 5 37 38.491 1
                                       3 3 33.579 0
```

Cross-validation creates many "analysis" and "assessment" sets from the same data set.



Review of R Programming Techniques

Putting commonly-used code in a function can speed coding and prevent copy/paste mistakes.

```
\verb| funct.name <- function(<arguments>){...code...}|\\
```

- Multiple commands are placed within braces { }
- Default values can be defined by setting function arguments equal to a value within the definition.
- The return keyword specifies the function return value.
 Otherwise the last value calculated in the function.
- The warning and stop functions can be used to throw warning and error messages.

```
sumn <- function(x, n=length(x)){sum(x[1:n], na.rm=TRUE)}
sumn(1:5, n=4)</pre>
[1] 10
```

If dealing with outside data/user input, it's good form to verify inputs. With your own code, it could help debug.

```
sumn <- function(x, n=length(x)){</pre>
  if(!is.numeric(x) || length(dim(x)) b <- c(1, 2, NA, 4, 5)
                                      sumn(a)
    stop("Requires one-dimensional
         numeric vector.")
  if(!is.numeric(n) || length(n) > 1) [1] 15
                                      sumn(a, n=3)
    stop("n must be a single
          numeric value.")
  if(round(n) != n){
                                      sumn(b, n=3)
   n <- round(n)</pre>
                                      [1] 3
    warning("n must be an integer
            and will be rounded.")
                                      sumn(a, n=3.2)
                                      Warning in sumn(a, n = 3.2): n must be
  if(n < 1 \mid | n > length(x))
                                                   and will be rounded.
    stop("n must be an integer between [1] 6
         1 and length of x.")
  return(sum(x[1:n], na.rm=TRUE))
```

Vectorized functions are usually more efficient than loops. Many R functions are vectorized.

```
Square roots of the whole
                                     Single values are repeated as
numbers from 1 to 5.
                                     needed in vectorized arithmetic.
a <- 1:5
                                     # Note: This is a long way to
                                     # calculate variance.
[1] 1 2 3 4 5
                                     а
                                     [1] 1 2 3 4 5
sqrt(a)
[1] 1.0000 1.4142 1.7321 2.0000 2.2361 a.mean <- mean(a)
                                     [1] 3
                                     a - a.mean
                                     [1] -2 -1 0 1 2
                                     sum((a - a.mean)^2) / (length(a) - 1)
                                     [1] 2.5
```

The s/lapply family of functions iterate functions over vector input. Usually more efficient than a loop.

- lapply always returns a list.
- sapply tries to return a simplified object.

The purrr package (in tidyverse) gives the map family of commands, which are similar to s/lapply.

```
# Note that the "suffix" gives the
map(1:5, sqrt)
                                       # desired type of output.
[[1]]
                                       map dbl(1:5, sqrt)
[1] 1
                                      [1] 1.0000 1.4142 1.7321 2.0000 2.2361
[[2]]
                                      # The "formula" format specifies
[1] 1.4142
                                      # an anonymous function compactly.
                                       \# using `.x` as the input.
[[3]]
                                       map_dbl(1:5, \sim sqrt(.x) + 2)
[1] 1.7321
                                       [1] 3.0000 3.4142 3.7321 4.0000 4.2361
[[4]]
[1] 2
[[5]]
[1] 2.2361
```

Sometimes a loop is just easier, especially if the function is complicated. So don't feel bad for using one if you have to!

```
for(i in 1:5){
                                     # To store loop results, it's good to
 # Note that `print` is needed
                                     # initialize variables first.
  # inside a loop to show results.
                                     a.sgrt <- numeric(5)
                                      # Note that these "type" commands with
 print(sqrt(i))
                                      # a number input give empty vectors.
                                     a.sqrt
[1] 1
[1] 1.4142
                                      [1] 0 0 0 0 0
[1] 1.7321
                                      for(i in 1:5){
                                       a.sqrt[i] <- sqrt(i)
[1] 2.2361
                                     a.sqrt
                                      [1] 1.0000 1.4142 1.7321 2.0000 2.2361
```