Lec 10: Classifiers

MATH 456 - Spring 2016

Navbar: [Home] [Schedule] [Data] [Week 13 Overview] [HW Info] [Google Group]

Assigned Reading & additional references

Reading: Afifi Chapter 11. Skim the math, read the rest carefully. Skip/skim: 11.9-11.11

Algorithm specific information is interspersed throughout the lecture notes.

- Lots of external reading/references
- Easy to go down the rabbit hole on this stuff!
- Read/skim around to better understand these methods, but know that we could have spent easily an entire semester on machine learning algorithms.
- This is just an introduction!
- Algorithms may take a while to computer, using cache=TRUE in the code chunk used to fit the model is advised.

Introduction to Classifiers

We will be discussing the following 6 classifying algorithms. These are also sometimes called *Machine Learning* (ML) algorithms. They are simply methods to build a model by learning from the relationships within the data.

- Logistic Regression
- Linear Discriminant analysis
- Naieve Bayes
- Decision Trees
- Random Forests
- k-nearest neighbors

For an idea of how many algorithms are out there:

- Data Mining Map: http://www.saedsayad.com/data_mining_map.htm
- http://topepo.github.io/caret/modelList.html

Most ML algorithms will use, in some way, every variable that you give it access to. To reduce model complexity and to reduce the chance of overfitting many models apply a *penalty* to the variable that diminishes their effect on the outcome, or reduces their chance of being included into the model.

Overfitting occurs when a statistical model describes random error or noise instead of the underlying relationship. Overfitting generally occurs when a model is excessively complex, such as having too many parameters relative to the number of observations. (Ref: https://en.wikipedia.org/wiki/Overfitting)

Machine learning in R: the caret package

We will be using the caret package in R to conduct all of our model building. Please read this small vignette for this package.

https://cran.r-project.org/web/packages/caret/vignettes/caret.pdf

Other packages you will need to install are: pROC, caTools, bnclassify, mda, party, rpart.plot, rattle, penalizedLDA.

- Typically don't need pre-install these
- You're prompted to install them when you use an algorithm that needs them
- Once installed, they will automatically load as needed.

If you want to use classifying algorithms other than the ones we have discussed here, these references will be useful:

- http://topepo.github.io/caret/bytag.html
- $\bullet \ \ http://artax.karlin.mff.cuni.cz/r-help/library/caret/html/train.html$

Cross-Validation

Many of these algorithms have tuning parameters that have to be optimized. The most common way to choose the optimal value of a tuning parameter is through cross validation (CV).

Recall that CV involves splitting the data (again) into testing and training data sets, running a single model over a range of values for the tuning parameter, and finding the value of the tuning parameter that provides the best fit.

We can specify how we want CV to occur in the trainControl function. Here we are specifying that we want to use repeated cross-validation. This means we apply k-fold CV (k defaults to 10 for this package) to one split of the data. Then repeat this process 2 more times on different rando testing/training splits of the data.

The classProbs=TRUE argument tells the function to calculate the probabilities of being in each class (depressed or not depressed), and to provide a summary information specific to two classes (confusion matrix).

Fitting (Training) the model

The workhorse in the caret package is the train() function. The generic syntax looks like the following:

```
train(y ~ . ,
    data=,
    method=,
    preProc = c("center", "scale"),
    metric = ,
    trControl = ctrl)
```

- y~. is the model syntax
- data is the data set used to train the model on
- method is the type of machine learning algorithm used

- preProc is the type of pre-processing that you want to be done
- metric is the performance metric that you want to optimize
- trControl is the type of training control you want to implement.

Unless otherwise specified, this is the training control method that we will use for all algorithms explored below.

Goal: Predict Depression

Note: The data management code file for the depression data set has been updated. Go download it and update your analysis data file.

```
depress <- read.delim("C:/GitHub/MATH456/data/depress_041616.txt")</pre>
```

Pre-processing the data

Assigned Reading: http://machinelearningmastery.com/how-to-prepare-data-for-machine-learning/

Often ML algorithms perform better when the variables are also centered and scaled. This can be done at the time of calling the algorithm, so we won't do that manually here.

Recoding and ensuring variables have correct attributes

In the data management code I have already recoded variables, ensured that all factor variables were being treated appropriately. This is also the step that you would conduct a PCA or create aggregate scales like CESD (which was already created for us).

However, specifically we need to ensure that R knows cases is a binary class variable with positive outcome (class 1) of "Depressed". The way I handle this is to first change cases to a factor variable, and swap the ordering of the levels. And then I apply named values to those levels.

```
depress$cases <- factor(depress$cases, levels=c(1,0))
levels(depress$cases) <- c("Depressed", "NotDepressed")</pre>
```

Identifying missing data

```
table(is.na(depress))

##
## FALSE TRUE
## 10876 2
```

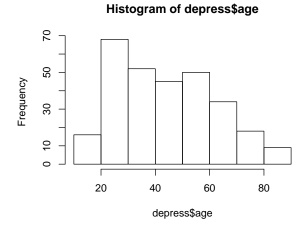
Two pieces of missing data. We have N=294, so will just delete those 2 records for now.

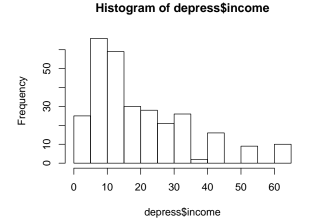
```
depress <- na.omit(depress)</pre>
```

Looking at distributions and outliers

Ensure each variable has sufficient variation.

```
par(mfrow=c(1,2))
hist(depress$age); hist(depress$income)
```





I then look at the categorical variables to see if there are any factor levels (categories) with a small number of records in them. This can lead to unstable estimates and algorithmic failure.

```
sapply(depress[,c('cases', 'marital', 'educat', 'employ', 'relig', 'health')], table)
```

```
## $cases
##
      Depressed NotDepressed
##
##
              49
                            243
##
   $marital
##
##
##
        Divorced
                          Married Never Married
                                                       Separated
                                                                         Widowed
##
                43
                              126
                                               73
                                                               12
                                                                               38
##
##
   $educat
##
##
             <HS
                             BS
                                      HS Grad
                                                                        PhD
                                                          MS
                             42
                                           114
                                                                          9
##
                                                          14
##
                       Some HS
   Some college
##
              48
                             61
##
   $employ
##
##
##
             FT Houseperson
                                 In School
                                                   Other
                                                                   PT
                                                                           Retired
##
            166
                           27
                                         2
                                                       4
                                                                    42
                                                                                 38
##
          Unemp
##
             13
```

```
##
## $relig
##
               3
                   4
##
     1
          2
##
   155
         51
             30
                  56
##
## $health
##
##
          2
               3
                   4
     1
                  14
## 129 114
             35
```

There are very few records with education less than HS, employed "in school" or "other". Specifically In school doesn't even have enough observations to calculate a variance on. I will combine "In school" with "Other".

```
library(car)
depress$employ <- recode(depress$employ, "'In School' = 'Other'")
table(depress$employ)

##
##
FT Houseperson Other PT Retired Unemp</pre>
```

As I was building these lecture notes, I also came across problems with the education variable (as expected). So I will further collapse categories for this variable.

42

38

13

6

##

Similarly I will look at the average value for all the 0/1 indicator variables in the data set to get an idea of the proportion of 1's. If any have a very low percent of events (1's) in the data we will have to be mindful of how the algorithms are performing with that variable in the model.

```
sapply(depress[,c('sex', 'drink', 'regdoc', 'treat', 'beddays', 'acuteill', 'chronill')], mean)
## sex drink regdoc treat beddays acuteill chronill
## 0.6198630 0.7945205 0.8150685 0.5068493 0.2157534 0.2979452 0.5068493
```

Manual variable selection.

Remove id, and the component variables c1:c20 that are used to create cesd, as well as the cesd variable since cases is a dichotomized version of this variable.

IMPORTANT NOTE At this point I have loaded some other package that has a **select** function. If you do not specify that here you want to specifically use the select function from the **dplyr** package, this WILL NOT WORK.

```
depress <- depress %>% dplyr::select(-id, -c1:-c20, -cesd)
names(depress)
```

```
## [1] "sex" "age" "marital" "educat" "employ" "income" ## [7] "relig" "cases" "drink" "health" "regdoc" "treat" ## [13] "beddays" "acuteill" "chronill"
```

The remaining variables are ones that I want to keep as candidate variables for a model.

Split the data into testing and training.

Instead of randomly sampling the entire data set to create the testing and training subsamples, the createDataPartition allows you to split the testing and training samples while stratifying on the outcome.

```
set.seed(1067)
inTrain <- createDataPartition(y=depress$cases, p=.7, list=FALSE)
train <- depress[inTrain,];dim(train)

## [1] 206   15

test <- depress[-inTrain,];dim(test)

## [1] 86   15</pre>
```

This is advantageous in that it ensures the relative proportion of the outcome variable is the same on both the testing and training data sets.

```
prop.table(table(depress$cases))
##
##
      Depressed NotDepressed
##
      0.1678082
                    0.8321918
prop.table(table(train$cases))
##
##
      Depressed NotDepressed
      0.1699029
                    0.8300971
##
prop.table(table(test$cases))
##
##
      Depressed NotDepressed
```

Now we are ready to build our models on the training data.

0.8372093

0.1627907

Use different classifying algorithms.

Logistic Regression

http://topepo.github.io/caret/Logistic_Regression.html

There are several different algorithms to perform a "flavor" of logistic regression analysis. We are going to use the LogitBoost algorithm.

Boosting methods have been originally proposed as ensemble methods, ... which rely on the principle of generating multiple predictions and majority voting (averaging) among the individual classifiers

Citation: https://web.stanford.edu/~hastie/Papers/buehlmann.pdf

Easy enough. Now let's look at the results of the model.

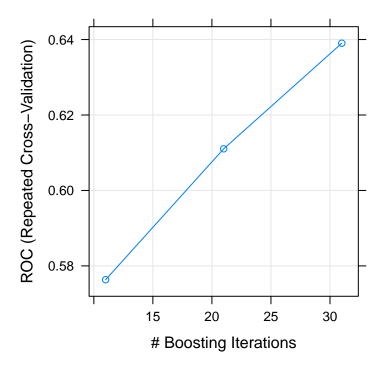
```
## Boosted Logistic Regression
##
## 206 samples
   14 predictor
##
    2 classes: 'Depressed', 'NotDepressed'
##
## Pre-processing: centered (22), scaled (22)
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 186, 186, 184, 185, 186, 185, ...
## Resampling results across tuning parameters:
##
##
    nIter ROC
                                             ROC SD
                                                        Sens SD
                                                                   Spec SD
                       Sens
                                  Spec
##
           0.5763344 0.222222
                                 0.9145969 0.1763731
                                                       0.2298731
                                                                   0.08139331
     11
##
    21
           0.6110430 0.2222222 0.8909586 0.1532994 0.2526993
                                                                   0.07477525
            0.6390251 0.2916667 0.8810458 0.1488553 0.2394818
##
##
## ROC was used to select the optimal model using the largest value.
## The final value used for the model was nIter = 31.
```

This output gives you a summary of the sample size going into the model, reminds you what the classes are, what pre-processing was done and what training control / optimization methods were used.

Then for each iteration the results show the sensitivity, specificity and area under the ROC curve.

We can also visualize how the AUC varied as a function of the number of boosting iterations (a tuning parameter).

```
plot(LogReg)
```



Almost all predictors available were used in the final model.

predictors(LogReg)

```
##
    [1] "income"
                              "beddays"
                                                    "age"
        "maritalWidowed"
                              "employUnemp"
##
    Γ41
                                                    "employPT"
                              "relig"
        "employHouseperson"
                                                    "sex"
        "employRetired"
                              "educatHS"
  [10]
                                                    "educatUG"
  [13]
        "health"
```

 $\begin{tabular}{ll} \bf Additional \ Reading \ on \ Penalized \ methods \ http://master.bioconductor.org/help/course-materials/2003/Milan/Lectures/anestisMilan3.pdf \end{tabular}$

Discriminant analysis

- Afifi Ch 11 introduces Linear Discriminant analyses
- No distributional assumptions
- Finds a combination of features that separates two events
- Not all combinations are linear. http://topepo.github.io/caret/Discriminant_Analysis.html

Linear Discriminant Analysis (LDA)

- Define Z to be a linear combination of X's: $Z = a_1X_1 + a_2X_2 + \dots + a_pX_p$
- Select a_i 's to maximize the Mahalanobis distance between the average value of Z in group 1 from the average value of Z in group 2.

Here I use two LDA algorithms: * LDA.1 uses the basic Linear Discriminant Analysis algorithm (method =lda) which does not contain any tuning parameters. * LDA.2 includes a stepwise feature selection procedure (method=stepLDA)

The stepwise LDA procedure is *very* verbose (it creates a lot of output) so I specifically am NOT showing the results here. (I put "'{r, results='hide'} in the code chunk header).

The results show that both have high sensitivity, low specificity, with AUC values between 0.5 and 0.7.

```
## Linear Discriminant Analysis
##
## 206 samples
   14 predictor
##
    2 classes: 'Depressed', 'NotDepressed'
##
##
## Pre-processing: centered (22), scaled (22)
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 185, 185, 185, 186, 185, 186, ...
## Resampling results
##
##
     ROC
               Sens
                          Spec
                                     ROC SD
                                                 Sens SD
                                                            Spec SD
##
     0.698602 0.1944444 0.9179739 0.1139862 0.2246041 0.07656073
##
##
## Linear Discriminant Analysis with Stepwise Feature Selection
##
## 206 samples
   14 predictor
     2 classes: 'Depressed', 'NotDepressed'
##
##
## Pre-processing: centered (22), scaled (22)
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 185, 186, 186, 185, 186, 185, ...
## Resampling results
##
##
                Sens
                            Spec
                                       ROC SD
                                                  Sens SD
                                                               Spec SD
##
     0.5002996 0.01944444
                            0.9727669 0.1322069
                                                  0.07480285 0.03991685
##
## Tuning parameter 'maxvar' was held constant at a value of Inf
##
```

```
## Tuning parameter 'direction' was held constant at a value of both
##
```

The finalModel item within the model object contains information on the prior probabilities of being depressed and the mean value for each variable within each of the depression categories. Specifically you can also output the values for the β regression coefficients. (Not all methods allow you to extract the values of the coefficients)

```
print(xtable(coef(LDA.1$finalModel), digits=2), type='latex')
```

	LD1
sex	-0.12
age	0.48
marital Married	-0.12
maritalNever Married	-0.12
maritalSeparated	-0.17
marital Widowed	-0.22
educatHS	-0.34
educatUG	-0.18
${\it employ} {\it House person}$	-0.27
employOther	-0.06
employPT	-0.36
employRetired	-0.02
employUnemp	-0.36
income	0.46
relig	-0.37
drink	-0.13
health	-0.03
regdoc	0.09
treat	-0.12
beddays	-0.36
acuteill	-0.06
chronill	-0.22

- Every variable in the data set was used in this model.
- Penalized models will *shrink* these estimates down to zero for variables that do not contribute.

Compare this to the LDA with a stepwise variable selection procedure added:

LDA.2finalModel

```
## method : lda
## final model : y ~ sex
## <environment: 0x000000000dace238>
##
## correctness rate = 0.83
```

This algorithm only chose gender as a predictor of depression.

• LDA compared to PCA: http://www.r-bloggers.com/computing-and-visualizing-lda-in-r/

Naieve Bayes

Bayesian probability models have the following form:

$$P(Y = 1|X) \propto \pi(Y = 1)P(X|Y = 1)$$

In english, this says that the posterior probability of an event occurring is proportional to the prior probability of the event occurring, times the likelihood of the event occurring given the observed data.

A naieve Bayes classifier can be thought of as fitting the joint probability model to optmize the joint likelihood p(Y, X) whereas logisic regression optimizes the conditional probability p(Y|X)

The Naieve Bayes classifier...

- is easy and fast, performs well with multiple class (categorical) predictions.
- Assumes the effect of x_1 on P(Y=1) is independent of the value of x_2 . (i.e. the X's are independent)
- Performs better than logistic regression with less training data if the assumption of conditional independence is upheld

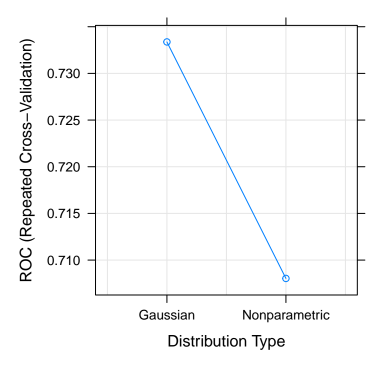
A list of Bayesian methods included in caret package. * http://topepo.github.io/caret/Bayesian_Model.html

This model takes slightly longer to fit compared to logistic regression, and at least in this example throws some warnings about zero probabilities

```
## Naive Bayes
##
## 206 samples
##
   14 predictor
##
    2 classes: 'Depressed', 'NotDepressed'
##
## Pre-processing: centered (22), scaled (22)
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 186, 185, 185, 186, 185, 185, ...
## Resampling results across tuning parameters:
##
##
     usekernel ROC
                           Sens
                                      Spec
                                                  ROC SD
                                                             Sens SD
##
    FALSE
                0.7333696 0.3972222 0.8379085
                                                 0.1128497
                                                             0.262324
      TRUE
                0.7080338  0.0000000  1.0000000
##
                                                 0.1305443 0.000000
##
     Spec SD
##
     0.07383915
     0.0000000
##
##
## Tuning parameter 'fL' was held constant at a value of 0
## ROC was used to select the optimal model using the largest value.
## The final values used for the model were fL = 0 and usekernel = FALSE.
```

This plot shows you the change in AUC as a function of if the algorithm was using a Gaussian or a non-parametric kernel.

plot(NB)



There is also a slightly different set of predictors chosen in the final model.

predictors(NB\$finalModel)

```
[1] "sex"
                                 "age"
                                                          "maritalMarried"
##
        "maritalNever.Married"
                                 "maritalSeparated"
                                                          "maritalWidowed"
##
        "educatHS"
                                 "educatUG"
                                                          "employHouseperson"
        "employOther"
                                 "employPT"
                                                          "employRetired"
##
   [10]
                                                          "relig"
        "employUnemp"
                                 "income"
                                                          "regdoc"
        "drink"
                                 "health"
        "treat"
                                                          "acuteill"
                                 "beddays"
   [19]
        "chronill"
```

Additional References * http://www.analyticsvidhya.com/blog/2015/09/naive-bayes-explained/ * http://www.saedsayad.com/naive_bayesian.htm

Decision Trees

Tree based methods are:

• Best when the relationship between the features and the response are complex

- lot of non-linear terms and interactions.
- Easy to explain & display graphically
- Mimics human decision making
- Tend to not have the same level of predictive accuracy as classical approaches.
- Non-parametric
- Can overfit the data more easily than other methods.

General Algorithm/Process

- 1. Find the variable split that best separates the categories of the response variable
- 2. Divide the data into two subsets based on that split
- 3. Within each subset, find the next variable that best separates the data into two categories.

The goal is to minimize the residual error. Technically trees can be built so tall that each observation is perfectly predicted. This is called *over fitting*. This can be avoided by *pruning* trees back (simplifying the tree).

- The level of pruning is a tuning parameter (complexity)
- Optimal complexity parameter can be determined by CV.

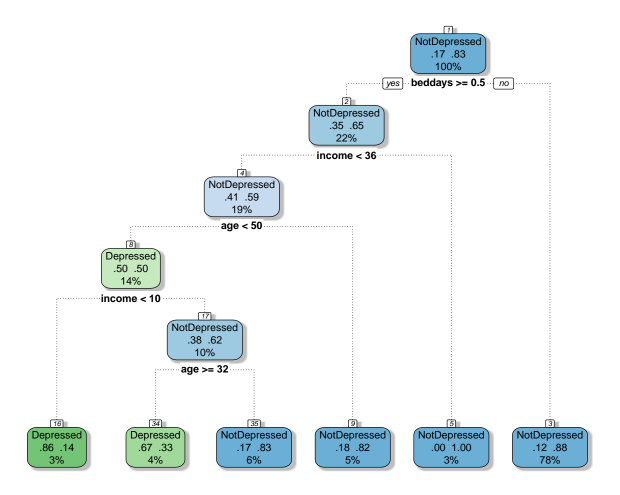
References & Readings

* https://en.wikipedia.org/wiki/Decision_tree_learning * https://rpubs.com/ryankelly/dtrees * http://trevorstephens.com/post/72923766261/titanic-getting-started-with-r-part-3-decision

```
## CART
##
## 206 samples
##
    14 predictor
     2 classes: 'Depressed', 'NotDepressed'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 186, 186, 186, 184, 185, 185, ...
## Resampling results across tuning parameters:
##
##
                 R.O.C
                                                       ROC SD
                                                                  Sens SD
                             Sens
                                           Spec
     ср
     0.0000000 0.5985566
##
                            0.086111111 0.9355120
                                                      0.1136176 0.15549226
##
     0.01785714 0.5916667 0.061111111 0.9531590
                                                      0.1193542 0.13118480
##
     0.03571429 \quad 0.5381536 \quad 0.008333333 \quad 0.9843137 \quad 0.1049947 \quad 0.04564355
##
     Spec SD
     0.07608103
##
     0.05627890
##
##
     0.04067599
##
## ROC was used to select the optimal model using the largest value.
## The final value used for the model was cp = 0.
```

Visualize the decision tree.

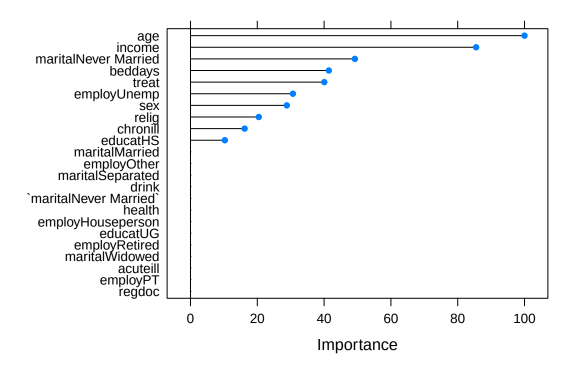
```
library(rpart.plot); library(rattle)
fancyRpartPlot(DT$finalModel, sub="")
```



The progrma can't start because libatk-1.0.0.dll is missing from your computer. Try reinstalling the program to fix the problem. Hit "OK" and install gTk2. You may have to restart R.

Tree based models also provide a measure on the importance of each candidate variable. These can be visualized using a **Variable Importance Plot**.

plot(varImp(DT))



This echo's the decision tree plot in that top three primary variables important in predicting depression is the number of days spent in bed, the level of general health, and employment status (specifically if they're unemployed)

Random Forests

explain this model

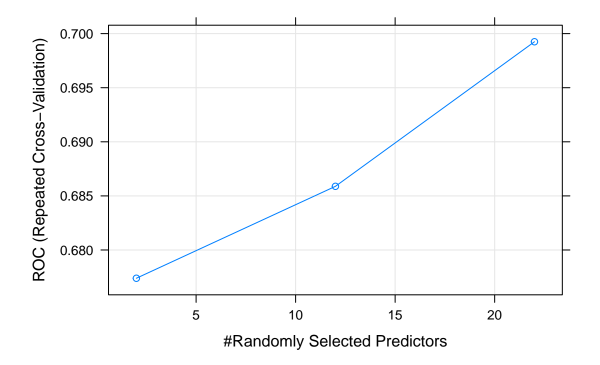
A random forest grows a large number of decision trees in order to improve classification rate.

http://trevorstephens.com/post/73770963794/titanic-getting-started-with-r-part-5-random/linear-part-5-random/lin

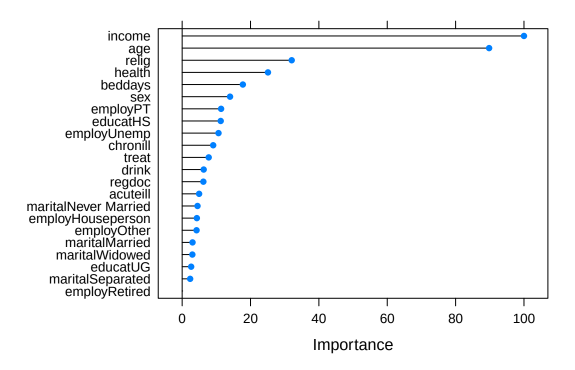
```
## Random Forest
##
## 206 samples
## 14 predictor
## 2 classes: 'Depressed', 'NotDepressed'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 186, 186, 185, 186, 185, ...
## Resampling results across tuning parameters:
```

```
##
                                                        Sens SD
##
     mtry ROC
                     Sens
                                  Spec
                                             ROC SD
           0.6773874 0.03055556
                                 1.0000000 0.1457678
                                                       0.09408598
##
##
     12
           0.6858842 0.12777778
                                 0.9745098
                                            0.1498292
                                                       0.15587679
     22
           0.6992466 0.11944444 0.9608932 0.1339350
##
##
     Spec SD
     0.00000000
##
     0.03993591
##
     0.04961685
##
##
## ROC was used to select the optimal model using the largest value.
## The final value used for the model was mtry = 22.
```

plot(RF)



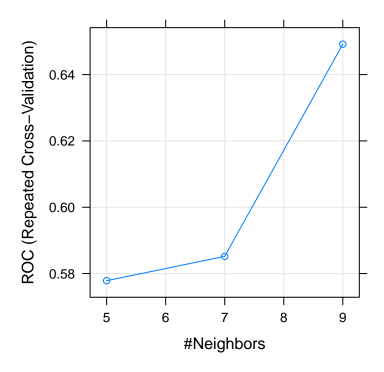
plot(varImp(RF))



k-nearest neighbors

explain this model

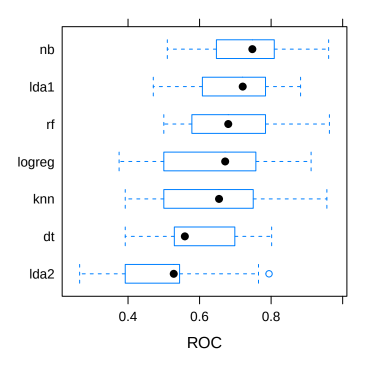
```
## k-Nearest Neighbors
## 206 samples
##
   14 predictor
##
     2 classes: 'Depressed', 'NotDepressed'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 185, 185, 185, 186, 186, ...
  Resampling results across tuning parameters:
##
##
    k ROC
                                          ROC SD
                                                     Sens SD
                                                                Spec SD
                   Sens
                               Spec
##
       0.5778459
                  0.09722222
                               0.9827887
                                          0.1509827
                                                     0.1419678
                                                                0.03378711
##
     7
       0.5851852 0.06944444
                               0.9789760
                                          0.1776012
                                                     0.1296239
                                                                0.04547457
##
       0.6491785 0.06944444
                              0.9924837
                                         0.1532432 0.1296239
                                                                0.03191753
##
## ROC was used to select the optimal model using the largest value.
## The final value used for the model was k = 9.
```



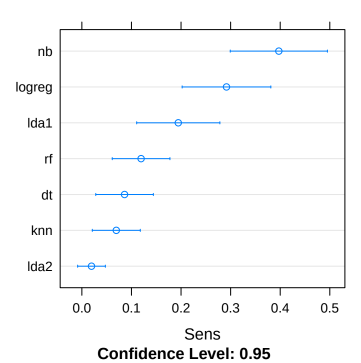
Compare algorithm performance

A resampling technique can be used to compare model performance. (Ref: _Hothorn at al, "The design and analysis of benchmark experiments- Journal of Computational and Graphical Statistics (2005) vol 14 (3) pp 675-699)_)

rValues <- resamples(list(logreg=LogReg,lda1=LDA.1, lda2=LDA.2, nb=NB, dt=DT, rf=RF, knn=kNN)) bwplot(rValues,metric="ROC")



dotplot(rValues,metric="Sens")



```
# Hint: Look at rValues$metrics for other metrics you can visualize & compare.
```

We can compare algorithms on their accuracy predicting the training data set. Here I create a confusion matrix by using each model to predict the class membership of data on the training data set.

```
acc.log <- confusionMatrix(predict(LogReg, train), train$cases, positive='Depressed')
acc.ld1 <- confusionMatrix(predict(LDA.1, train), train$cases, positive='Depressed')
acc.ld2 <- confusionMatrix(predict(LDA.2, train), train$cases, positive='Depressed')
acc.nb <- confusionMatrix(predict(NB, train), train$cases, positive='Depressed')
acc.dt <- confusionMatrix(predict(DT, train), train$cases, positive='Depressed')
acc.rf <- confusionMatrix(predict(RF, train), train$cases, positive='Depressed')
acc.knn <- confusionMatrix(predict(kNN, train), train$cases, positive='Depressed')</pre>
```

Ok so how do extract the accuracy values? Lets see what data is in the object created by the confusionMatrix function.

```
names(acc.log)
## [1] "positive" "table"
                              "overall" "byClass"
                                                    "dots"
names(acc.log$overall)
## [1] "Accuracy"
                         "Kappa"
                                          "AccuracyLower"
                                                           "AccuracyUpper"
                         "AccuracyPValue" "McnemarPValue"
## [5] "AccuracyNull"
names(acc.log$byClass)
## [1] "Sensitivity"
                               "Specificity"
                                                      "Pos Pred Value"
## [4] "Neg Pred Value"
                               "Prevalence"
                                                      "Detection Rate"
## [7] "Detection Prevalence" "Balanced Accuracy"
```

Now that I've identified that the value for Accuracy is the first element in the **\$overall** list, I can extract these values and compare them in a tabular format. Furthermore the sensitivity, specificity and other table values can be found in the **\$byClass** list.

For berivity sake (and to demonstrate some advanced R programming techniques) I create a function getmetrics to extract only the values of the performance metrics I am interested in, then apply that function to all models in a list.

	Accuracy	Sensitivity	Specificity
Boosted Logistic Regression	0.883	0.486	0.965
LDA	0.874	0.429	0.965
LDA w/stepwise	0.830	0.000	1.000
Naieve Bayes	0.791	0.429	0.865
Decision Tree	0.869	0.343	0.977
Random Forest	1.000	1.000	1.000
k-Nearest neighbors	0.845	0.086	1.000

Making predictions

On the hold-out testing data set.

```
confusionMatrix(predict(RF, test), test$cases, positive='Depressed')
```

```
## Confusion Matrix and Statistics
##
##
                 Reference
## Prediction
                  Depressed NotDepressed
                           2
##
     Depressed
                                        3
                         12
##
     NotDepressed
                                       69
##
##
                  Accuracy : 0.8256
##
                    95% CI : (0.7287, 0.899)
       No Information Rate: 0.8372
##
       P-Value [Acc > NIR] : 0.67881
##
##
##
                     Kappa: 0.1365
   Mcnemar's Test P-Value : 0.03887
##
##
##
               Sensitivity: 0.14286
##
               Specificity: 0.95833
##
            Pos Pred Value: 0.40000
##
            Neg Pred Value: 0.85185
##
                Prevalence: 0.16279
##
            Detection Rate: 0.02326
##
      Detection Prevalence: 0.05814
##
         Balanced Accuracy: 0.55060
##
##
          'Positive' Class : Depressed
##
```

The Random Forest model had an accuracy of 0.837(0.742,0.908) on the testing data set.

Additional references

- http://www.statmethods.net/advstats/discriminant.html
- $\bullet \ \ http://rstudio-pubs-static.s3.amazonaws.com/35817_2552e05f1d4e4db8ba87b334101a43da.html$
- https://www.youtube.com/watch?v=s8pvp2Ctxfc
- http://www.r-bloggers.com/in-depth-introduction-to-machine-learning-in-15-hours-of-expert-videos/

 $http://michael.hahsler.net/SMU/EMIS7332/R/chap5.html \\ [top]$

On Your Own

On Your Own

Using the Parental HIV data set, build a predictive model for whether or not a student skips class <code>HOOKEY</code> using all 6 of the classifying algorithms discussed in this set of lecture notes.