# Decision Trees

This script will go over Decision Trees in R. It will not cover generalization such as SEM Trees or Rasch Trees

#### Resources:

Basic intro to Decision Trees: http://www.statmethods.net/advstats/cart.html Full list of data mining packages in R: http://cran.r-project.org/web/views/MachineLearning.html

For longitudinal data:

REEMtree

Two packages will be used and their caret equivalents:

rpart (tree accomplishes very similar thing):http://cran.r-project.org/web/packages/rpart/vignettes/longintro.pdf

party: http://cran.r-project.org/web/packages/party/vignettes/party.pdf

```
In caret, method =
"rpart" - tuning = cp (complexity parameter)
"rpart2" - tuning = maxdepth
"rpartCost" - tuning = cp and cost
"ctree" - tuning = mincriterion (p value thresholds)
"ctree2" - tuning = maxdepth
(see "train_model_list" in caret reference manual)
```

### Bonus:

non-greedy tree algorithm:

evtree: http://cran.r-project.org/web/packages/evtree/vignettes/evtree.pdf

Lets load the main packages

```
library(caret)
library(pROC)
library(randomForest)
library(ada)
library(ISLR)
library(party)
library(MASS) # for boston data
data(Boston)
```

# Regression (continous outcome)

Use rpart first with the Boston data use regression first – predicting median value of homes

```
#str(Boston)

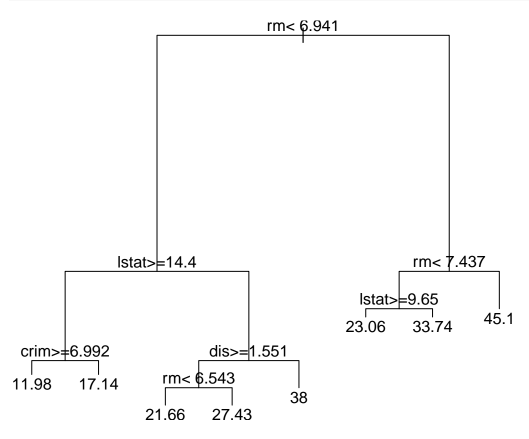
# lets get a baseline with linear regression
lm.Boston <- lm(medv ~., data=Boston)
#summary(lm.Boston)</pre>
```

We do pretty well with linear regression R-squared of .74

# **CART**

How about if we just blindly apply Decision Trees

```
rpart.Boston <- rpart(medv ~., data=Boston)
#summary(rpart.Boston)
plot(rpart.Boston); text(rpart.Boston)</pre>
```



# this can be hard to interpret, so I like to look at a different output rpart.Boston

```
## n = 506
##
## node), split, n, deviance, yval
         * denotes terminal node
##
##
    1) root 506 42716.3000 22.53281
##
##
      2) rm< 6.941 430 17317.3200 19.93372
##
        4) lstat>=14.4 175 3373.2510 14.95600
          8) crim>=6.99237 74 1085.9050 11.97838 *
##
##
          9) crim< 6.99237 101 1150.5370 17.13762 *
##
        5) lstat< 14.4 255 6632.2170 23.34980
##
         10) dis>=1.5511 248 3658.3930 22.93629
##
           20) rm< 6.543 193 1589.8140 21.65648 *
```

```
##
            21) rm>=6.543 55 643.1691 27.42727 *
##
          11) dis< 1.5511 7 1429.0200 38.00000 *
##
       3) rm>=6.941 76 6059.4190 37.23816
         6) rm< 7.437 46 1899.6120 32.11304
##
##
          12) lstat>=9.65 7 432.9971 23.05714 *
          13) lstat< 9.65 39 789.5123 33.73846 *
##
##
         7) rm>=7.437 30 1098.8500 45.09667 *
pred1 <- predict(rpart.Boston)</pre>
cor(pred1,Boston$medv)**2
## [1] 0.8075721
Doing really well – Rsquared = 0.81
## Lasso Regression
What if we tried regularized (penalized) regression instead?
Note: for glmnet, both the x's and y have to be in separate matrices
- and all class = numeric
- don't worry about response, doesn't have to be factor for logistic
— just specify "binomial"
y.B <- Boston$medv
x.B <- sapply(Boston[,-14],as.numeric)</pre>
```

```
y.B <- Boston$medv
x.B <- sapply(Boston[,-14],as.numeric)

# alpha =1 for lasso, 0 for ridge
library(glmnet)
cv <- cv.glmnet(x.B,y.B,alpha=1)
lasso.reg <- glmnet(x.B,y.B,alpha=1,family="gaussian",lambda=cv$lambda.min)

lasso.resp <- predict(lasso.reg,newx=x.B)
cor(y.B,lasso.resp)**2</pre>
```

```
## s0
## [1,] 0.7400946
```

Taking into account cross-validation, we do worse compared to linear regression with no tuning.

So the plot for rpart didn'tcome out that well. Good news, there are better options for plotting.

 $http://blog.revolution analytics.com/2013/06/plotting-classification- and -regression-trees-with-plotrpart. \\ html$ 

Let's load some new packages:

```
library(rattle)
```

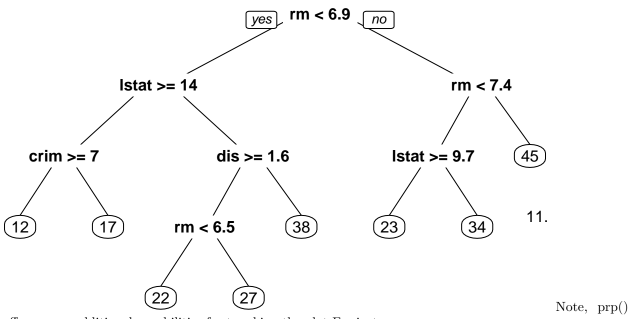
```
## Rattle: A free graphical interface for data mining with R.
## Version 3.3.0 Copyright (c) 2006-2014 Togaware Pty Ltd.
## Type 'rattle()' to shake, rattle, and roll your data.
```

```
library(rpart.plot)
library(RColorBrewer)
library(partykit)
```

Note: rattle is package that uses a GUI (think SPSS) for data mining applications check out book: http://www.amazon.com/Data-Mining-Rattle-Excavating-Knowledge/dp/1441998896

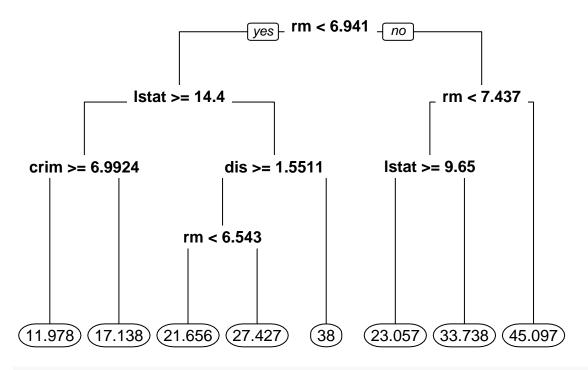
Anyways, lets try some new, prettier plots:

```
# prp(); from rpart.plot
prp(rpart.Boston); text(rpart.Boston)
```

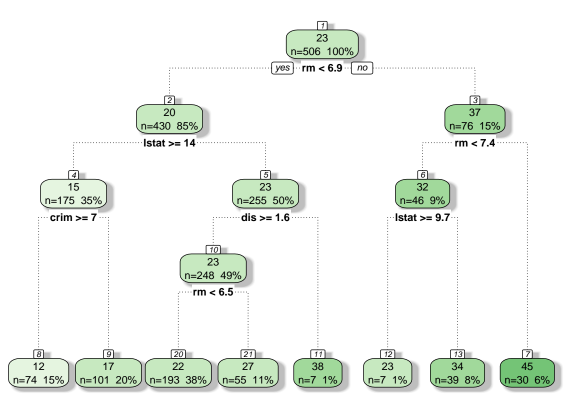


offers many additional capabilities for tweaking the plot For instance:

```
# ?prp
prp(rpart.Boston, varlen=10, digits=5, fallen.leaves=T)
```



#fancyRpartPlot(); from rattle
fancyRpartPlot(rpart.Boston)



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## **Conditional Inference Trees**

So what about with conditional inference trees?

What if we want a smaller tree? This can be accomplished a number of ways. We can prespecify the maxdepth, the minimum number of people per node, as well as making more restrictive splitting criterion.

Example of prespecifying the depth with ctree()

```
ctree.Boston <- ctree(medv ~., data=Boston)
#plot(ctree.Boston) # too big of a tree
pred2 <- predict(ctree.Boston)
cor(pred2,Boston$medv)**2</pre>
```

```
## [1] 0.8746338
```

We do better than rpart, Rsquared = 0.87

Biggest difference between ctree() and rpart() is that ctree() does not demonstrate bias with respect to the number of response options, and supposedly had less of a propensity to overfit than rpart(). Note: the models are not optimizing based on Rsquared, most likely MSE

So what do we think now? Are we happy with results? Remember, decision trees are generally quite robust, so it may not be necessary to check assumptions. – See Table 10.1 ESL

But what about generalizability?

Although not as serious as with SVM for instance, Decision Trees have a propensity to overfit, meaning the tree structure won't generalize well

So let's try just creating a simple Training and Test datasets

```
train = sample(dim(Boston)[1], dim(Boston)[1]/2) # half of sample
Boston.train = Boston[train, ]
Boston.test = Boston[-train, ]
```

Try linear regression first

```
lm.train <- lm(medv ~., data=Boston.train)
pred.lmTest <- predict(lm.train,Boston.test)
cor(pred.lmTest,Boston.test$medv)**2</pre>
```

```
## [1] 0.7255828
```

Note: we are taking our lm object trained on the train dataset, and using these fixed coefficients to predict values on the test dataset.

In SEM, this is referred to as a tight replication strategy No difference in using a test dataset – both Rsq are 0.74

How about with rpart?

```
rpart.train <- rpart(medv ~., data=Boston.train)

pred.rpartTest <- predict(rpart.train,Boston.test)
cor(pred.rpartTest,Boston.test$medv)**2</pre>
```

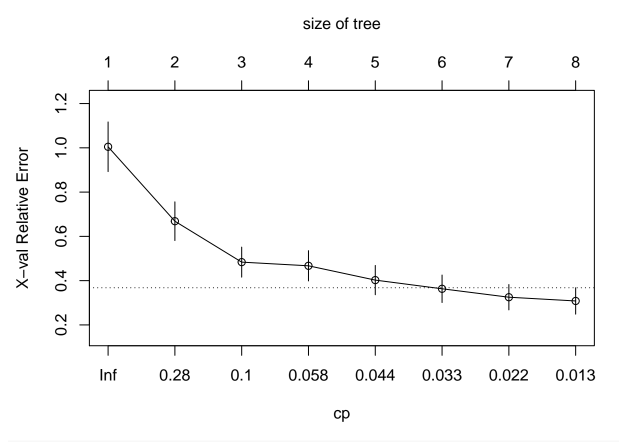
## ## [1] 0.7727604

Not as good – drops from 0.81 to 0.76 – still better than lm() But with rpart, it is common to prune trees back. What if we try this, is there less of a drop in  $\mathbb{R}^2$ ?

Note: rpart automatically does internal CV, varying the complexity paramter (cp). If you use the tree package instead, you will have to use cv.tree()

With plotcp() we are going to choose the error within 1 SE of the lowest cross-validated error. This will be used to prune

# plotcp(rpart.train)



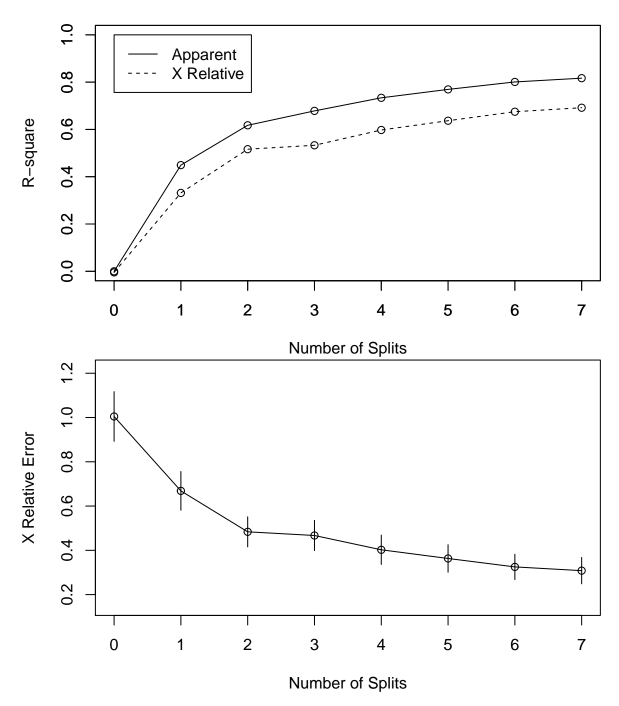
## printcp(rpart.train)

```
##
## Regression tree:
## rpart(formula = medv ~ ., data = Boston.train)
##
## Variables actually used in tree construction:
## [1] crim dis lstat rm
```

```
##
## Root node error: 21116/253 = 83.46
##
## n= 253
##
##
          CP nsplit rel error xerror
## 1 0.449032 0 1.00000 1.00472 0.112126
## 2 0.168432
                 1 0.55097 0.66853 0.087437
## 3 0.060925
                2 0.38254 0.48363 0.067807
## 4 0.055291
                 3 0.32161 0.46707 0.068421
## 5 0.035603
                 4 0.26632 0.40244 0.066157
## 6 0.031519
                5 0.23072 0.36319 0.062115
## 7 0.015921
                 6 0.19920 0.32513 0.057282
## 8 0.010000
                7 0.18328 0.30822 0.059545
```

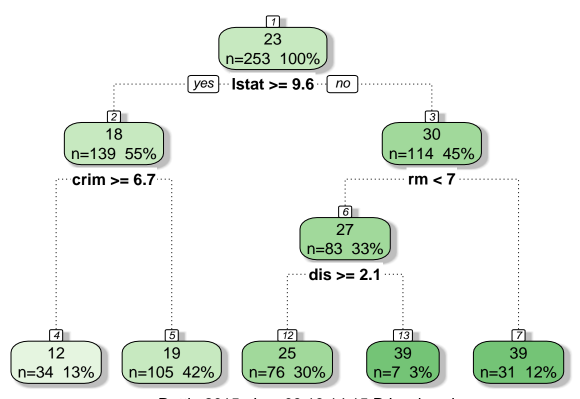
### rsq.rpart(rpart.train)

```
## Regression tree:
## rpart(formula = medv ~ ., data = Boston.train)
## Variables actually used in tree construction:
## [1] crim dis
                lstat rm
##
## Root node error: 21116/253 = 83.46
## n= 253
##
          CP nsplit rel error xerror
##
## 1 0.449032
                  0 1.00000 1.00472 0.112126
## 2 0.168432
                  1
                      0.55097 0.66853 0.087437
## 3 0.060925
                  2 0.38254 0.48363 0.067807
## 4 0.055291
                3 0.32161 0.46707 0.068421
## 5 0.035603
                 4 0.26632 0.40244 0.066157
## 6 0.031519
                 5 0.23072 0.36319 0.062115
## 7 0.015921
                6 0.19920 0.32513 0.057282
             7 0.18328 0.30822 0.059545
## 8 0.010000
```



```
prune.Bos <- prune(rpart.train,0.053)

#plot(prune.Bos); text(prune.Bos)
fancyRpartPlot(prune.Bos)</pre>
```



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```
pred.prune <- predict(prune.Bos,Boston.test)
cor(pred.prune,Boston.test$medv)</pre>
```

## [1] 0.8342364

caret:

```
ctree.train <- ctree(medv ~., data=Boston.train)
#plot(ctree.train) # huge tree
pred.ctreeTest <- predict(ctree.train,Boston.test)
cor(pred.ctreeTest,Boston.test$medv)**2</pre>
```

## [1] 0.7593072

It is worth noting how much more of an effect there was for using a test dataset with the tree methods as compared to lm(), this is pretty typical, and much more important with more "flexible" methods such as random forests, gbm, svm etc...

# Classification (categorical outcome)

### Two Biggest Things To Remember:

- 1. Make sure functions outcome variable is categorical; as.factor(outcome)
- 2. Using predict() changes. Variable across packages

As a baseline, we will use logistic regression.

```
library(ISLR)
data(Default)
head(Default)
```

```
##
     default student
                       balance
                                  income
## 1
         Nο
                  No 729.5265 44361.625
## 2
          No
                 Yes 817.1804 12106.135
## 3
                  No 1073.5492 31767.139
          Nο
## 4
          No
                  No
                      529.2506 35704.494
## 5
          No
                     785.6559 38463.496
## 6
          No
                 Yes 919.5885 7491.559
```

#### str(Default)

```
## 'data.frame': 10000 obs. of 4 variables:
## $ default: Factor w/ 2 levels "No","Yes": 1 1 1 1 1 1 1 1 1 1 1 1 1 ...
## $ student: Factor w/ 2 levels "No","Yes": 1 2 1 1 1 2 1 2 1 1 ...
## $ balance: num 730 817 1074 529 786 ...
## $ income : num 44362 12106 31767 35704 38463 ...
```

My favorite function in R is str(), as it gives the class of each variable and other summary characteristics. Most important thing to note is that the "default" variable is already coded as a factor variable, meaning that R now knows it is categorical, and will change the cost function (thus estimator) accordingly.

This is really important because rpart,randomForest and other packages do not automatically detect whether it is a regression or classification problem. If you don't change the outcome variable to its proper class, you could get a suboptimal answer (use the wrong estimator i.e. regression instead of logistic regression)

Now let's do logistic regression

```
lr.out <- glm(default~student+balance+income,family="binomial",data=Default)
summary(lr.out)</pre>
```

```
##
## Call:
  glm(formula = default ~ student + balance + income, family = "binomial",
      data = Default)
##
##
## Deviance Residuals:
      Min
                1Q
                     Median
                                  3Q
                                          Max
## -2.4691 -0.1418 -0.0557 -0.0203
                                       3.7383
##
## Coefficients:
                Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) -1.087e+01 4.923e-01 -22.080 < 2e-16 ***
## studentYes -6.468e-01 2.363e-01
                                    -2.738
                                            0.00619 **
## balance
               5.737e-03 2.319e-04
                                     24.738
                                             < 2e-16 ***
## income
               3.033e-06 8.203e-06
                                      0.370 0.71152
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
```

```
##
## Null deviance: 2920.6 on 9999 degrees of freedom
## Residual deviance: 1571.5 on 9996 degrees of freedom
## AIC: 1579.5
##
## Number of Fisher Scoring iterations: 8
```

I always find it much harder to figure out how well I am doing with logistic regression. One of the best ways to assess results in my opinion is the use of receiver operating characteristic curves (ROC curves).

good intro to using ROC:

https://ccrma.stanford.edu/workshops/mir2009/references/ROCintro.pdf

These plots are a balanace of sensitivity and specificity. Ideally the curve gets as close as possible to the upper left corner.

To get this plot, we need to get our predictions from our logistic model.

```
glm.probs=predict(lr.out,type="response")
#glm.pred00=ifelse(glm.probs>0.5,1,0)

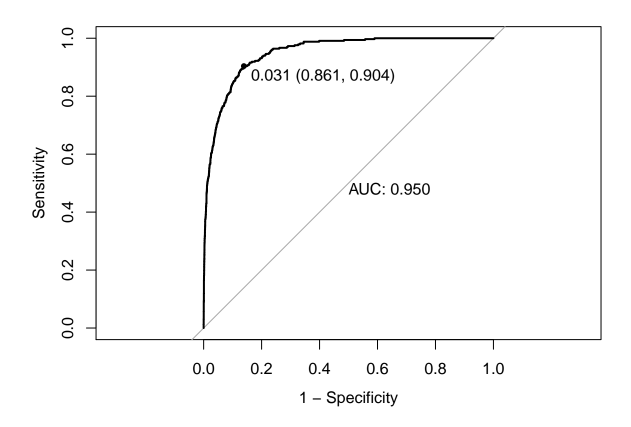
rocCurve <- roc(Default$default,glm.probs)
pROC::auc(rocCurve)

## Area under the curve: 0.9496

pROC::ci(rocCurve)

## 95% CI: 0.9402-0.959 (DeLong)

# quartz()
plot(rocCurve, legacy.axes = TRUE,print.thres=T,print.auc=T)</pre>
```



```
##
## Call:
## roc.default(response = Default$default, predictor = glm.probs)
##
## Data: glm.probs in 9667 controls (Default$default No) < 333 cases (Default$default Yes).
## Area under the curve: 0.9496</pre>
```

For AUC (area under the curve), values of 0.8 and 0.9 are good (the higher the better)

## predict() with missing variables

How about lasso logistic regression?

```
library(glmnet)
yy = as.numeric(Default$default)
xx = sapply(Default[,2:4],as.numeric)
lasso.out <- cv.glmnet(xx,yy,family="binomial",alpha=1,nfolds=10) #alpha=1 == lasso; 0 =
# find best lambda
ll <- lasso.out$lambda.min

lasso.probs <- predict(lasso.out,newx=xx,s=ll,type="response")</pre>
```

Results from lasso using CV

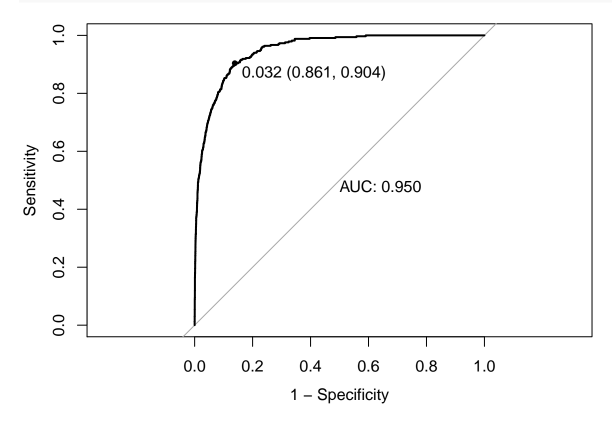
```
rocCurve.lasso <- roc(Default$default,lasso.probs)
pROC::auc(rocCurve.lasso)</pre>
```

```
## Area under the curve: 0.9495
```

```
pROC::ci(rocCurve.lasso)
```

```
## 95% CI: 0.9401-0.959 (DeLong)
```

```
# quartz()
plot(rocCurve.lasso, legacy.axes = TRUE,print.thres=T,print.auc=T)
```



```
##
## Call:
## roc.default(response = Default$default, predictor = lasso.probs)
##
## Data: lasso.probs in 9667 controls (Default$default No) < 333 cases (Default$default Yes).
## Area under the curve: 0.9495</pre>
```

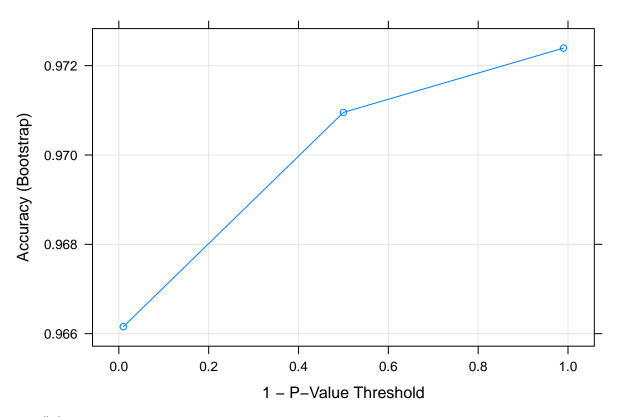
Almost identical results to logistic regression with no penalization.

# Using Decision Trees for Classification

Instead of demonstrating how to use rpart() or ctree(), I prefer to use the train() from caret. This makes it much easier to test out multiple different methods, as well as automatically vary the tuning parameters such as depth, complexity etc..

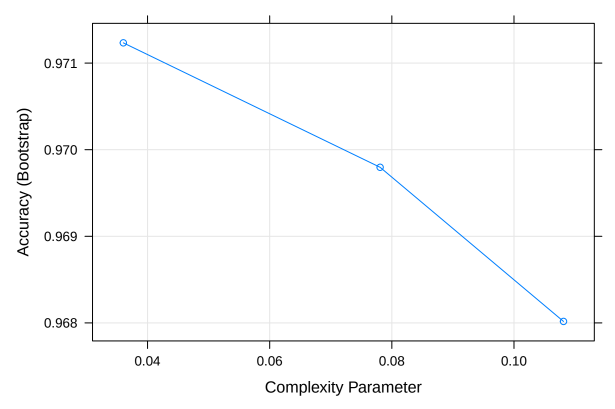
train() for ctree

train.ctree <- train(as.factor(default)~student+balance+income,data=Default,method="ctree")
plot(train.ctree)</pre>



train() for rpart

train.rpart <- train(as.factor(default)~student+balance+income,data=Default,method="rpart")
plot(train.rpart)</pre>



In train() and through trainControl() you can see that it automatically varies different tuning parameters (see caret documentation for the different options for each method), while defaulting to bootstrap estimation to test out each. This is a great way to prevent overfitting.

In examining both plots, it seems as both methods do comparably well, while also they both have different tuning parameters (X-axis). Based on these plots, I would increase the number of values for the tuning parameters, as the accuracy did not reach a maximum necessarily outside of the tails. (tuneLength = 3 is default)

### using a confusion matrix

```
train.class <- predict(train.rpart,Default,type="raw")
confusionMatrix(train.class,Default$default)</pre>
```

```
## Confusion Matrix and Statistics
##
##
             Reference
##
  Prediction
                No
                     Yes
                     243
##
          No
              9639
##
          Yes
                28
                      90
##
##
                  Accuracy : 0.9729
##
                     95% CI: (0.9695, 0.976)
##
       No Information Rate: 0.9667
##
       P-Value [Acc > NIR] : 0.0002082
##
##
                      Kappa: 0.3885
    Mcnemar's Test P-Value : < 2.2e-16
##
```

```
##
##
               Sensitivity: 0.9971
##
               Specificity: 0.2703
            Pos Pred Value: 0.9754
##
##
            Neg Pred Value: 0.7627
                Prevalence: 0.9667
##
##
            Detection Rate: 0.9639
      Detection Prevalence: 0.9882
##
##
         Balanced Accuracy: 0.6337
##
##
          'Positive' Class : No
##
# much better than just using
table(train.class,Default$default)
##
## train.class
                 No
                     Yes
##
           No 9639
                     243
                      90
##
           Yes
                 28
```

# Changing the cutoff for class assignment

This uses the optimal cutoff from the pROC plot

```
train.prob <- predict(train.rpart,Default,type="prob")[,2]
train.class2 <- ifelse(train.prob > .031,1,0)
confusionMatrix(as.numeric(Default$default)-1,train.class2,positive="1")
```

```
## Confusion Matrix and Statistics
##
##
            Reference
## Prediction
                0
##
            0 9541 126
            1 171 162
##
##
##
                  Accuracy: 0.9703
                    95% CI: (0.9668, 0.9735)
##
##
      No Information Rate: 0.9712
##
       P-Value [Acc > NIR] : 0.71715
##
##
                     Kappa: 0.5065
   Mcnemar's Test P-Value: 0.01068
##
##
##
               Sensitivity: 0.5625
##
               Specificity: 0.9824
##
            Pos Pred Value: 0.4865
##
            Neg Pred Value: 0.9870
##
                Prevalence: 0.0288
##
            Detection Rate: 0.0162
##
     Detection Prevalence: 0.0333
##
         Balanced Accuracy: 0.7724
```

```
##
## 'Positive' Class : 1
##

# much better than just using
table(train.class2, Default$default)

##
## train.class2 No Yes
## 0 9541 171
## 1 126 162
```

If there is a large class imbalance, may want to try unbalance sampling: http://www.win-vector.com/blog/2015/02/does-balancing-classes-improve-classifier-performance/

Another way to get optimal threshold:

# Boosting and Random Forests

For this, we will use the **caret** package as an interface to both the **gbm** and **randomForest** packages.

Because gbm and random forests take much longer to run, we could set up parallelization through caret and other packages. http://topepo.github.io/caret/parallel.html

In my experience, unless your dataset is huge, parallelization with random forests tends to take longer than setting up only serial computation.

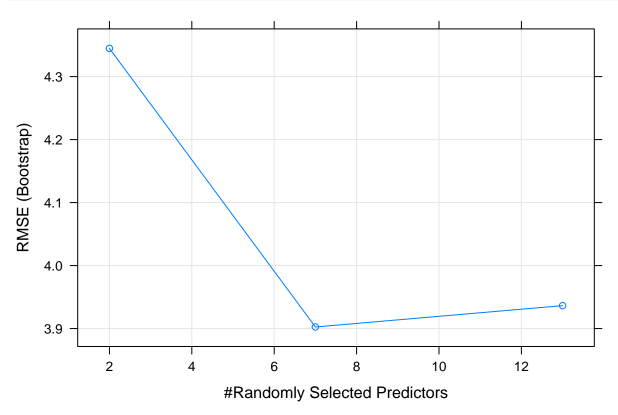
In caret, randomForest has two implementations, method="rf" and method="parRF" with parRF being the parallel version. The only tuning parameter for both is mtry.

Note, that using the train() will take longer, as it is using different tuning parameters and by default using bootstrap sampling to prevent overfitting.

To let train() pick the values of mtry, just set tuneLength to however many different values you want it to try. Default is 3.

```
## Random Forest
##
## 253 samples
    13 predictor
##
##
## No pre-processing
## Resampling: Bootstrapped (25 reps)
## Summary of sample sizes: 253, 253, 253, 253, 253, 253, ...
##
## Resampling results across tuning parameters:
##
##
           RMSE
                     Rsquared
                                RMSE SD
                                            Rsquared SD
     mtry
##
      2
           4.344815
                     0.7934294
                                0.4972010
                                            0.05267354
##
      7
           3.902491
                     0.8211233
                                0.4824882
                                            0.04325310
##
     13
           3.936368
                     0.8140024
                                0.4915482
                                            0.04527352
##
## RMSE was used to select the optimal model using the smallest value.
## The final value used for the model was mtry = 7.
```

## plot(train.rf1)



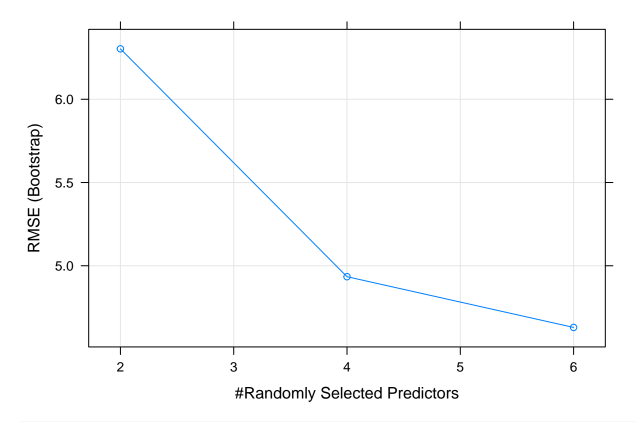
```
#rf <- train.rf1$finalModel
varImp(train.rf1)</pre>
```

```
## rf variable importance
##
## Overall
```

```
## rm
           100.000
## lstat
           95.158
            37.869
## dis
            37.432
## crim
## tax
            29.496
## nox
            26.076
## ptratio 24.686
## indus
            21.866
## age
            21.053
            6.600
## black
## rad
             4.364
             2.117
## zn
             0.000
## chas
# see how we do on hold out sample
pred.test1 <- predict(train.rf1,Boston.test)</pre>
cor(pred.test1,Boston.test$medv)**2
## [1] 0.8882108
cforest is implemented as method="cforest" with the only tuning parameter being mtry
train.cf1 <- train(medv ~., data=Boston.train,method="cforest")</pre>
train.cf1
## Conditional Inference Random Forest
##
## 253 samples
## 13 predictor
##
## No pre-processing
## Resampling: Bootstrapped (25 reps)
##
## Summary of sample sizes: 253, 253, 253, 253, 253, 253, ...
## Resampling results across tuning parameters:
##
##
     mtry RMSE
                     Rsquared
                                RMSE SD
                                            Rsquared SD
##
           6.302850 0.6377645 0.6536649
                                            0.12335194
##
           4.934225 0.7274101 0.7533223 0.08943965
##
           4.629622 0.7412545 0.7030520 0.07741343
##
## RMSE was used to select the optimal model using the smallest value.
```

plot(train.cf1)

## The final value used for the model was mtry = 6.



# varImp(train.cf1)

```
## cforest variable importance
##
##
            Overall
## lstat
            100.0000
## rm
            47.8977
## crim
             8.0861
             4.4242
## ptratio
## dis
             3.9706
## tax
             3.5950
## indus
             3.5160
## black
             2.3331
             2.1976
## nox
## age
             2.0309
             1.2058
## rad
## chas
             0.7659
## zn
             0.0000
```

```
# see how we do on hold out sample
pred.test2 <- predict(train.cf1,Boston.test)
cor(pred.test2,Boston.test$medv)**2</pre>
```

# ## [1] 0.8583804

For classification, a couple other things to use paired with train()

```
# set up data
data(Carseats)
#attach(Carseats)
#hist(Carseats$Sales)
High=ifelse(Carseats$Sales<=8,"No","Yes")
Carseats=data.frame(Carseats, High)
Carseats$Sales <- NULL
Carseats$ShelveLoc <- as.numeric(Carseats$ShelveLoc)

train2 = sample(dim(Carseats)[1], dim(Carseats)[1]/2) # half of sample
Carseats.train = Carseats[train2, ]
Carseats.test = Carseats[-train2, ]</pre>
```

Now run random forests

```
train.rf2 <- train(as.factor(High) ~ ., data=Carseats.train,method="rf",trControl=cont)</pre>
train.rf2
rf.probs=predict(train.rf2,newdata=Carseats.test,type="prob")[,2]
rocCurve22 <- roc(Carseats.train$High,rf.probs)</pre>
auc(rocCurve22)
plot(rocCurve22)
rf.class=predict(train.rf2,newdata=Carseats.train)
confusionMatrix(Carseats.train$High,rf.class)
# how about on test dataset
rf.classTest=predict(train.rf2,Carseats.test)
confusionMatrix(Carseats.test$High,rf.classTest)
# ROC
rf.probs1=predict(train.rf2, Carseats.test, type="prob")[,2]
rocCurve4 <- roc(Carseats.test$High,rf.probs1)</pre>
auc(rocCurve4)
ci(rocCurve4)
plot(rocCurve4)
```

cforest for binary – automatically knows it is binary, unlike randomForest

!! bug with the predict.cforest function

```
train.cf2 <- train(High ~ ., data=Carseats.train,method="cforest",trControl=cont)
train.cf2

cf.probs=predict(train.cf2) # doesn't work. ????
rocCurve33 <- roc(Carseats$High,cf.probs)
auc(rocCurve33)
plot(rocCurve33)</pre>
```

```
cf.class=predict(train.cf2$finalModel)
confusionMatrix(Carseats$High,cf.class) # no errors

# how about on test dataset

cf.classTest=predict(train.cf2,newdata=Carseats.test)
confusionMatrix(Carseats.test$High,cf.classTest)
```

## boosting

packages: "gbm" and "ada" – both can be accessed through caret

Example tuning parameters for "gbm: http://topepo.github.io/caret/training.html

For this, I am going to use method="ada" which is one of the forms of boosting. Currently, problem with method="gbm"

Basic set up to compare results to RF:

```
#modelLookup("ada")
train.ada <- train(as.factor(High) ~ ., data=Carseats.train,method="ada",trControl=cont)
train.ada

ada.probs=predict(train.ada,newdata=Carseats.test,type="prob")[,2]
rocCurve3 <- roc(Carseats.test$High,ada.probs)
auc(rocCurve3)
ci(rocCurve3)
plot(rocCurve3,add=T,col=c(2)) # color red is ada</pre>
```