Predictive Analytics: practical 2 solutions

```
library("caret")
data(FuelEconomy, package = "AppliedPredictiveModeling")
set.seed(25)
```

Cross validation and the bootstrap

• Fit a linear regression model to the cars2010 data set with FE as the response, using EngDispl, NumCyl and NumGears as predictors.

```
the response, using EngDispl, NumCyl and NumGears as predictors. The data set can be loaded data("FuelEconomy", package = mLM = train(FE ~ EngDispl + NumCyl + NumGears, method = "lm", data = "ApplicativeModeling").
```

• What is the training error rate (RMSE) for this model?

```
res = resid(mLM)
(trainRMSE = sqrt(mean(res * res)))
## [1] 4.59
```

Hint: The training error can be found by taking the square root of the average square residuals. The sqrt and resid functions may be useful.

• Re-train your model using the validation set approach to estimate a test RMSE, make your validation set equivalent to a half of the full available data.

• How does this compare to the training error that we estimated above?

```
# it's larger, often training error under estimates test error
getTrainPerf(mLMVS)

## TrainRMSE TrainRsquared method
## 1    4.847    0.616    lm

trainRMSE

## [1] 4.59
```

• Go through the same process using the different methods for estimating test error. That is leave one out and *k*–fold crossvalidation as well as bootstrapping.

```
# set up train control objects
tcLOOCV = trainControl(method = "LOOCV")
tcKFOLD = trainControl(method = "cv", number = 10)
tcBOOT = trainControl(method = "boot")
```

10-fold cross validation can be shown to be a good choice for almost any situation.

```
# train the model
mLML00CV = train(FE ~ EngDispl + NumCyl + NumGears, method = "lm", data = cars2010,
    trControl = tcL00CV)
mLMKFOLD = train(FE ~ EngDispl + NumCyl + NumGears, method = "lm", data = cars2010,
   trControl = tcKF0LD)
mLMB00T = train(FE ~ EngDispl + NumCyl + NumGears, method = "lm", data = cars2010,
trControl = tcBOOT)
```

• How do these estimates compare with the validation set approach?

```
getTrainPerf(mLMVS)
## TrainRMSE TrainRsquared method
## 1 4.847 0.616 lm
getTrainPerf(mLML00CV)
## TrainRMSE TrainRsquared method
## 1 4.612 0.6214 lm
getTrainPerf(mLMKF0LD)
## TrainRMSE TrainRsquared method
## 1 4.585 0.6315 lm
getTrainPerf(mLMB00T)
## TrainRMSE TrainRsquared method
## 1 4.567 0.6326 lm
# all lower than validation set, we mentioned it tended to over estimate
```

• The object returned by train also contains timing information that can be accessed via the times component of the list. Which of the methods is fastest?

The \$ notation can be used pick a single list component.

```
mLMVS$times$everything
   user system elapsed
  0.244 0.000 0.247
mLML00CV$times$everything
   user system elapsed
  4.796 0.008 4.823
mLMKF0LD$times$everything
   user system elapsed
  0.280 0.004 0.283
mLMB00T$times$everything
##
   user system elapsed
## 0.344 0.004 0.350
```

• Using k-fold cross validation to estimate test error investigate how the number of folds effects the resultant estimates and computation

```
# a number of trainControl objects
tc2 = trainControl(method = "cv", number = 2)
tc5 = trainControl(method = "cv", number = 5)
tc10 = trainControl(method = "cv", number = 10)
tc15 = trainControl(method = "cv", number = 15)
tc20 = trainControl(method = "cv", number = 20)
# train the model using each
mLM2 = train(FE ~ EngDispl + NumCyl + NumGears, method = "lm", data = cars2010,
        trControl = tc2)
mLM5 = train(FE ~ EngDispl + NumCyl + NumGears, method = "lm", data = cars2010,
        trControl = tc5)
mLM10 = train(FE ~ EngDispl + NumCyl + NumGears, method = "lm", data = cars2010,
        trControl = tc10)
mLM15 = train(FE ~ EngDispl + NumCyl + NumGears, method = "lm", data = cars2010,
        trControl = tc15)
mLM20 = train(FE ~ EngDispl + NumCyl + NumGears, method = "lm", data = cars2010,
        trControl = tc20)
# use a data frame to store all of the relevant information
(info = data.frame(Folds = c(2, 5, 10, 15, 20), Time = c(mLM2$times$everything[1],
         \verb|mLM5$ times everything[1]|, \verb|mLM10$ times everything[1]|, \verb|mLM15$ times everything[1]|,
         mLM20$times$everything[1]), Estimate = c(mLM2$results$RMSE, mLM5$results$RMSE,
         mLM10$results$RMSE, mLM15$results$RMSE, mLM20$results$RMSE)))
## Folds Time Estimate
## 1
                2 0.264 4.595
## 2
                  5 0.260 4.597
## 3 10 0.280 4.604
## 4 15 0.300 4.583
## 5
             20 0.316 4.551
# as there are more folds it takes longer to compute, not an issue with such
# a small model but something to consider on more complicated models.
# Estimates are going down as the number of folds increases. This is
# because for each held out fold we are using a greater proportion of the
# data in training so expect to get a better model.
```

• Experiment with adding terms to the model, transformations of the predictors and interactions say and use cross validation to estimate test error for each. What is the best model you can find? You can still use the validate and mark functions to look at how your models fair on the unseen data.

Penalised regression

The diabetes data set in the lars package contains measurements of a number of predictors to model a response y, a measure of disease progression. There are other columns in the data set which contain interactions so we will extract just the predictors and the response. The data has already been normalized.

```
## load the data in
data(diabetes, package = "lars")
diabetesdata = cbind(diabetes$x, y = diabetes$y)
```

 Try fitting a lasso, ridge and elastic net model using all of the main effects, pairwise interactions and square terms from each of the predictors.1

```
model formula = as.formula(paste("y~(.)^2 + ", paste0("I(", colnames(drabetes data) ctor a polynomial term)) colnames (drabetes data) ctor a polynomial term) collaborated col
                              "^2)", collapse = "+")))
mLASSO = train(modelformula, data = diabetesdata, method = "lasso")
 mRIDGE = train(modelformula, data = diabetesdata, method = "ridge")
mENET = train(modelformula, data = diabetesdata, method = "enet")
```

 Try to narrow in on the region of lowest RMSE for each model, don't forget about the tuneGrid argument to the train function.

best is

mLASSOfiner\$bestTune

```
<sup>1</sup> Hint: see notes for shortcut on creating
model formula. Also be aware that if the
doesn't make sense
```

fraction = 0 is the same as the null model.

 $y \sim (.) \land 2$ is short hand for a model that includes pairwise interactions for each predictor, so if we use this we

```
# examine previous output then train over a finer grid near the betteshootld only need to add the square
mLASSOfine = train(modelformula, data = diabetesdata, method = "lassderm%uneGrid = data.frame(fraction = seq(0.1
   0.5, by = 0.05)))
mLASSOfine$results
   fraction RMSE Rsquared RMSESD RsquaredSD
## 1
       0.10 16.88  0.9531  1.198  0.005822
## 2
        0.15 17.06 0.9513 1.371 0.006878
## 3
        0.20 17.36  0.9496  1.456  0.007611
## 4
       0.25 17.47 0.9490 1.422 0.007500
## 5
       0.30 17.56 0.9485 1.415 0.007553
## 6
        0.35 17.63 0.9481 1.423 0.007681
        0.40 17.70 0.9477 1.418
## 7
                                  0.007710
## 8
        0.45 17.75
                   0.9474 1.413
                                  0.007728
## 9
        0.50 17.80 0.9472 1.413
                                  0.007802
# best still right down at the 0.1 end
mLASSOfiner = train(modelformula, data = diabetesdata, method = "lasso", tuneGrid = data.frame(fraction = seq(0.
   0.15, by = 0.01))
mLASSOfiner$results
##
     fraction RMSE Rsquared RMSESD RsquaredSD
## 1
        ## 2
         0.02 38.43 0.9546 18.834 0.004405
## 3
        0.03 30.59 0.9545 17.109 0.005125
        0.04 25.97 0.9540 14.843
## 4
                                   0.005017
## 5
                    0.9540 12.779
        0.05 23.24
                                   0.004980
## 6
        0.06 21.58 0.9532 10.932
                                   0.005182
## 7
        0.07 20.60 0.9527 9.168 0.005150
## 8
        0.08 19.90 0.9522 7.476 0.005188
## 9
        0.09 19.28 0.9517 5.914 0.005419
## 10
         0.10 18.72 0.9513 4.526
                                  0.005481
         0.11 18.25
                    0.9509 3.331
                                   0.005565
## 11
## 12
         0.12 17.88
                    0.9505 2.334
                                   0.005592
## 13
         0.13 17.62
                    0.9502 1.615
                                   0.005573
## 14
        0.14 17.45 0.9501 1.218
                                   0.005595
## 15
        0.15 17.37 0.9499 1.123
                                    0.005772
```

```
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## fraction
## 15 0.15
mRIDGEfine = train(modelformula, data = diabetesdata, method = "ridge", tuneGrid = data.frame(lambda = seq(0,
   0.1, by = 0.01))
mRIDGEfine$results
##
   lambda RMSE Rsquared RMSESD RsquaredSD
## 1 0.00 18.54 0.9436 1.5934 0.010896
## 2
     0.01 17.16 0.9514 0.7828 0.005701
## 3
     0.02 17.07 0.9519 0.7510 0.005412
      0.03 17.09 0.9518 0.7575 0.005467
## 4
## 5
      0.04 17.18 0.9512 0.7813
                               0.005677
     0.05 17.31 0.9505 0.8134 0.005966
## 6
## 7
     0.06 17.49 0.9494 0.8494 0.006297
## 8
     0.07 17.69 0.9483 0.8867 0.006651
## 9
     0.08 17.91 0.9470 0.9239 0.007016
## 10 0.09 18.16 0.9456 0.9602 0.007385
mRIDGEfiner = train(modelformula, data = diabetesdata, method = "ridge", tuneGrid = data.frame(lambda = seq(0.06)
   0.03, by = 0.001))
mRIDGEfiner$results
   lambda RMSE Rsquared RMSESD RsquaredSD
## 1 0.005 16.98 0.9524 1.055 0.005344
## 2 0.006 16.95 0.9526 1.061 0.005314
    0.007 16.92 0.9527 1.067
## 3
                               0.005295
## 4
     0.008 16.90 0.9528 1.073
                               0.005285
## 5 0.009 16.89 0.9529 1.079 0.005282
## 6 0.010 16.88 0.9529 1.084 0.005282
## 7 0.011 16.87 0.9530 1.090 0.005287
## 8 0.012 16.86 0.9530 1.095 0.005294
## 9 0.013 16.86 0.9530 1.101 0.005304
## 10 0.014 16.86 0.9530 1.106 0.005316
## 11 0.015 16.86 0.9530 1.111
                               0.005330
## 12  0.016 16.86  0.9530  1.116
                               0.005345
## 13 0.017 16.86 0.9530 1.121 0.005362
## 14 0.018 16.87 0.9529 1.126 0.005380
## 15 0.019 16.87 0.9529 1.131 0.005399
## 16 0.020 16.88 0.9529 1.136 0.005420
## 17 0.021 16.89 0.9528 1.140 0.005441
## 18 0.022 16.90 0.9528 1.145 0.005464
## 19 0.023 16.90
                 0.9527 1.150
                               0.005487
## 20 0.024 16.91 0.9527 1.154
                               0.005511
## 21 0.025 16.92 0.9526 1.159 0.005536
## 22 0.026 16.94 0.9525 1.163 0.005561
## 23 0.027 16.95 0.9525 1.168 0.005587
## 24 0.028 16.96 0.9524 1.172 0.005614
## 25 0.029 16.97 0.9523 1.176 0.005641
## 26 0.030 16.99 0.9522 1.181 0.005669
# the best one
```

```
##
   lambda
## 10 0.014
```

mRIDGEfiner\$bestTune

```
mENETfine = train(modelformula, data = diabetesdata, method = "enet", tuneGrid = expand.grid(lambda = c(0.001,
  0.01, 0.1), fraction = c(0.4, 0.5, 0.6))
mENETfine$results
## lambda fraction RMSE Rsquared RMSESD RsquaredSD
## 1 0.001 0.4 16.26 0.9565 0.8848 0.003724
## 4 0.010
             0.4 16.18  0.9581 1.0260  0.003675
## 7 0.100 0.4 22.94 0.9553 2.1321 0.004127
## 2 0.001
            0.5 16.64 0.9546 0.8795 0.003647
## 5 0.010 0.5 15.89 0.9584 0.8946 0.003591
## 8 0.100
             0.5 17.05 0.9559 0.9928 0.003703
## 3 0.001
             0.6 16.84 0.9535 0.9578 0.004200
             0.6 16.23 0.9566 0.8379 0.003280
## 6 0.010
           0.6 16.74 0.9536 0.8435 0.004589
## 9 0.100
mENETfiner = train(modelformula, data = diabetesdata, method = "enet", tuneGrid = expand.grid(lambda = seq(0.001
   0.1, length.out = 10), fraction = 0.5))
mENETfiner$results
   lambda fraction RMSE Rsquared RMSESD RsquaredSD
## 1 0.001 0.5 16.85 0.9535 0.8808 0.005661
## 2 0.012
              0.5 15.98 0.9578 0.7835 0.003957
## 3 0.023 0.5 16.03 0.9576 0.8601 0.003964
## 4 0.034
             0.5 16.18  0.9570 0.9269  0.004037
## 5 0.045
             0.5 16.36  0.9566 0.9742  0.004042
             0.5 16.53 0.9562 1.0125 0.003977
## 6 0.056
## 7
      0.067
             0.5 16.70
                         0.9558 1.0340 0.003958
## 8 0.078 0.5 16.84 0.9555 1.0587
                                       0.004075
## 9 0.089 0.5 16.96 0.9552 1.0920 0.004301
## 10 0.100 0.5 17.06 0.9549 1.1178 0.004488
mENETfiner$bestTune
## fraction lambda
## 2 0.5 0.012
```

We can view the coefficients via

```
coef = predict(mLASSO$finalModel,
 mode = "fraction",
 s = mLASSO$bestTune$fraction,# which ever fraction was chosen as best
 type = "coefficients"
```

 How many features have been chosen by the lasso and enet models?

```
# use predict to find the coefficients
coefLASSO = predict(mLASSOfiner$finalModel, mode = "fraction", type = "coefficient",
   s = mLASSO$bestTune$fraction, )
sum(coefLASS0$coefficients != 0)
## [1] 57
coefENET = predict(mENETfiner$finalModel, mode = "fraction", type = "coefficient",
  s = mENET$bestTune$fraction)
sum(coefENET$coefficients != 0)
## [1] 24
```

• How do these models compare to principal components and partial least squares regression?

```
mPCR = train(modelformula, data = diabetesdata, method = "pcr", tuneGrid = data.frame(ncomp = 1:7))
mPLS = train(modelformula, data = diabetesdata, method = "pls", tuneGrid = data.frame(ncomp = 1:7))
mPLS2 = train(modelformula, data = diabetesdata, method = "pls", tuneGrid = data.frame(ncomp = 5:15))
getTrainPerf(mLASSOfiner)
## TrainRMSE TrainRsquared method
## 1 17.37 0.9499 lasso
getTrainPerf(mRIDGEfiner)
## TrainRMSE TrainRsquared method
## 1 16.86
                   0.953 ridge
getTrainPerf(mENETfiner)
## TrainRMSE TrainRsquared method
## 1 15.98 0.9578 enet
getTrainPerf(mPCR)
## TrainRMSE TrainRsquared method
## 1 16.53 0.9557 pcr
getTrainPerf(mPLS2)
## TrainRMSE TrainRsquared method
## 1 16.05 0.9587 pls
# The elastic net model has the lowest estimated test error, all are fairly
# similar. The elastic net model suggests only 21 non--zero coefficients out
# of all of those included in the model.
```