Modeling with k-means, Part 1

This project shows how we can use the k-means algorithm to build better classification models than what construct using various supervised learning approaches. Here I am employing k-means as a supervised giving it information about the labels indirectly in two distinct ways: (1) through input from one or more su algorithms in the form of an additional variable, or probability column; and (2) through the addition of weights arrived at through cross-validation scores that rely on the labels, or res

Even when k-means is used for unsupervised learning, there are a number of ways we guide, or constrai For example, (a) we tell it the number of clusters to find; (b) we select the variables we want the algorithn (c) we choose the transformations to apply to those variables. Variable selection and variable transformathe results we see when applying k-means. The less appropriate the inputs to the algorithm are, the less will be.

A fourth way we can guide the algorithm is by applying weights to the variables. While this is a variant of own category because the weights I refer to appear in the object dissimilarity measure---the distance forr weights are used to regulate a variable's relative influence "in determining the overall dissimilarity betwee observations]" (p.505 of *The Elements of Statistical Learning, Second Edition*, by Hastie, Tibshirani, and Equation (14.24)). In the modeling that follows, the weight given to a variable depends on the role the var presence of the other variables, in partitioning the data into the subgroups that are of interest to us. Thos assigned a greater weight are, we might say, more important for correctly identifying these subgroups, wl assigned a smaller weight are less important. But even if we can identify the more important predictors ir very likely we will not know what weights to assign unless we have data to work with in which the subgroup already identified, or labeled. When k-means is used for unsupervised learning, this information is not available.

Sometimes, however, we do have the data we need in order to assign meaningful and relevant weights. I scenarios that are of interest to me in this project. In other words, we can employ k-means as a supervise algorithm. But rather than directly giving it the labels to train on, we can give it the probability output from classification algorithms. Under certain conditions, this hybrid approach yields an even better classifier. A this project is to identify these conditions.

In Part1 I apply this hybrid approach to data on downer cows. Here the k-means hybrid model outperforn classification models surveyed, and weights can be used to improve the model. In Part2 I apply the hybri California housing dataset. Here optimal weights are much harder to establish (the number of predictors doubles, and the amount of training data is 33X greater). Weights in this example do not help to improve much.

In Part 4 I work with the Italian wine dataset used by Kabacoff (R in Action, p.370); there are 3 varieties to

* * * * *

The downer cow data for Part 1 are introduced on p.266 of Sanford Weisberg's Applied Linear Regressic

"For unknown reasons, dairy cows sometimes become recumbent---they lay down. Called *downers*, thes a serious illness that may lead to death of the cow. These data are from a study of blood samples of over studied at the Ruakura New Zealand Animal Health Laboratory during 1983-1984. A variety of blood test and for many of the animals, the outcome (survived, died, or animal was killed) was determined. The goa survival can be predicted from the blood measurements. The variables in the data file *downer.txt* are desermined data were collected from veterinary records, and not all variables were recorded for all cows."

```
In [ ]: require(car)
        require(repr)
        require(ggplot2)
        require(stringr)
        require(faraway)
        require(parallel)
        require(randomForest)
        require(gbm)
        require(plyr)
        require(e1071)
In [2]: options(digits= 5, show.signif.stars= FALSE)
In [4]: | df <- read.table("/home/greg/Documents/stat/alr3data/downer.txt", header= TRL</pre>
        dim(df)
        colnames(df)
         435 · 9
         'Calving' · 'Daysrec' · 'CK' · 'AST' · 'Urea' · 'PCV' · 'Inflamat' · 'Myopathy' · 'Outcome'
In [ ]: # Description of the variables in the dataset:
        # Outcome : = 1 if the cow survived; 0 if it died or was killed
        ### Categorical variables:
        # Calving : = 0 if measurements were taken before calving; = 1 if taken after
        # Inflamat : Is inflammation present? 0 = no; 1 = yes.
        # Myopathy : Is muscle disorder present? 0 = no; 1 = yes.
        ### Continuous variables:
        # Daysrec : days recumbent when measurements were taken.
        # CK : serum creatine phosphokinase (in U/l at 30C)
        # AST : serum asparate amino transferase (in U/l at 30C)
        # Urea : serum ureas (in mmol/l)
        # PCV : packed cell volume (hematocrit), percent
In [4]: # 38% of the 435 cows survived.
        table(df$Outcome)
        round(mean(df$Outcome, na.rm=TRUE), 4)
```

760 166

```
In [5]: table(df$Calving, useNA= "ifany")
                 1 <NA>
            0
          107 324
 In [6]: table(df$Myopathy, useNA= "ifany")
                 1 <NA>
          127
                95 213
 In [7]: summary(df$Daysrec)
            Min. 1st Qu.
                          Median
                                    Mean 3rd Qu.
                                                     Max.
                                                             NA's
           0.000
                   0.000
                           1.000
                                    1.947 3.000 20.000
 In [3]: # Percent of cows recumbent < 1 day.</pre>
         val <- dim(df[which(df$Daysrec < 1),])[1]</pre>
         round(val/nrow(df), 3)
         0.308
 In [8]: summary(df$CK)
            Min. 1st Qu.
                          Median
                                    Mean 3rd Qu.
                                                     Max.
                                                             NA's
              13
                     560
                            1760
                                     5352
                                             5467
                                                    71000
 In [9]: summary(df$AST)
            Min. 1st Qu.
                          Median
                                    Mean 3rd Qu.
                                                             NA's
                                                     Max.
            33.0
                  123.0
                           240.0
                                    398.4 492.0
                                                  2533.0
                                                                6
In [10]: summary(df$Urea)
                                                             NA's
            Min. 1st Qu.
                          Median
                                    Mean 3rd Qu.
                                                     Max.
           1.000
                   5.625
                           7.600
                                    9.803 10.975 50.000
                                                              169
In [11]: summary(df$PCV)
            Min. 1st Qu.
                                    Mean 3rd Qu.
                                                             NA's
                          Median
                                                     Max.
           13.00 32.00
                           35.00
                                    35.56 40.00
                                                    61.00
                                                              260
 In [1]: ### COMMENTS:
         # There is a great deal of missing data for PCV, Urea, Myopathy,
         # and Inflamat. Because of this, I do not want to include any
         # of these variables in my classifiers. I want to keep as many
         # of the 435 records as possible since I will be comparing the
         # performance of different classifiers, and the greater the
         # number of observations I have to work with, the smaller the
         # variability in these performance measures.
```

```
b0 <- as.numeric(coeffs[1])</pre>
             b1 <- as.numeric(coeffs[2])</pre>
             prob <- (\exp(b0 + b1*x)/(1 + \exp(b0 + b1*x)))
             return(round(prob, 3))
In [15]: # To what degree does Calving "explain" Outcome?
         fit_calv <- glm(Outcome ~ Calving, data=df, family= binomial())</pre>
         summary(fit calv)
         Call:
         glm(formula = Outcome ~ Calving, family = binomial(), data = df)
         Deviance Residuals:
                                     30
            Min
                    1Q Median
                                            Max
          -1.01
                  -1.01
                         -0.89
                                   1.35
                                            1.50
         Coefficients:
                     Estimate Std. Error z value Pr(>|z|)
                                   0.206 -3.50 0.00046
         (Intercept)
                     -0.721
         Calving
                        0.321
                                   0.235
                                            1.36 0.17228
         (Dispersion parameter for binomial family taken to be 1)
             Null deviance: 573.6 on 430 degrees of freedom
         Residual deviance: 571.7 on 429 degrees of freedom
           (4 observations deleted due to missingness)
         AIC: 575.7
         Number of Fisher Scoring iterations: 4
In [32]: # Is Daysrec a predictor of Outcome?
         fit_daysre <- glm(Outcome ~ Daysrec, data=df, family= binomial())</pre>
         summary(fit_daysre)
         Call:
         glm(formula = Outcome ~ Daysrec, family = binomial(), data = df)
         Deviance Residuals:
                    1Q Median
                                     30
                                           2.288
         -1.103 -1.034 -0.842 1.254
         Coefficients:
                     Estimate Std. Error z value Pr(>|z|)
                                  0.1308
         (Intercept) -0.1782
                                           -1.36 0.17311
         Daysrec
                      -0.1689
                                  0.0512
                                           -3.30 0.00097
         (Dispersion parameter for binomial family taken to be 1)
             Null deviance: 574.57 on 431 degrees of freedom
         Residual deviance: 561.62 on 430 degrees of freedom
           (3 observations deleted due to missingness)
         AIC: 565.6
```

```
In [34]: # The probability that the average cow survives increases as
         # the number of days recumbent decreases.
         get prob(0, fit daysre$coeff)
         0.456
 In [ ]: ### COMMENT:
         # Since Daysrec is predictive of the outcome, we perhaps
         # ought to control for this variable when regressing
         # Outcome on the blood measurements.
         # Further downstream I regress CK and AST on Daysrec.
         # The results tell us that we should control for
         # Daysrec.
In [31]: # Is Calving predictive when we control for Daysrec?
         fit_daysre_calv <- glm(Outcome ~ Daysrec + Calving, data=df,</pre>
                                  family= binomial())
         summary(fit_daysre_calv)
         Call:
         glm(formula = Outcome ~ Daysrec + Calving, family = binomial(),
              data = df
         Deviance Residuals:
                     1Q Median
                                       30
                                              Max
          -1.146 -1.026 -0.834 1.230
                                            2.468
         Coefficients:
                      Estimate Std. Error z value Pr(>|z|)
          (Intercept)
                       -0.3682
                                    0.2268
                                             -1.62 0.10447
          Daysrec
                       -0.1877
                                    0.0537
                                              -3.49 0.00048
                                    0.2398
                                              1.22 0.22158
         Calving
                        0.2931
          (Dispersion parameter for binomial family taken to be 1)
              Null deviance: 569.75 on 427 degrees of freedom
         Residual deviance: 553.09 on 425 degrees of freedom
            (7 observations deleted due to missingness)
          AIC: 559.1
         Number of Fisher Scoring iterations: 4
 In [3]: # Function for computing "proportion of deviance explained", an
         # R-sqrd statistic for link= logit. See p.41 of Julian Faraway's # "Extending the Linear Model with R" (2006: Chapman & Hall).
         # This statistic is due to N. Nagelkerke (1991; see Faraway's
         # bibliography).
         get_RsqrdDev <- function(modl) {</pre>
              n obs <- length(modl$fitted.values)</pre>
              ans <- (1-exp((modl$deviance - modl$null.deviance)/n obs))/(1-exp(-modl$r
```

```
summary(fit_ck)
        glm(formula = Outcome \sim I(log(CK)), family = binomial(), data = df)
        Deviance Residuals:
                    10 Median
                                   30
                                          Max
                                1.059
         -2.134 -0.881 -0.561
                                        1.994
        Coefficients:
                    Estimate Std. Error z value Pr(>|z|)
                                0.5809
         (Intercept)
                    4.0006
                                        6.89 5.7e-12
         I(log(CK))
                     -0.6117
                                0.0793
                                         -7.71 1.2e-14
         (Dispersion parameter for binomial family taken to be 1)
            Null deviance: 550.49 on 412 degrees of freedom
        Residual deviance: 475.18 on 411 degrees of freedom
           (22 observations deleted due to missingness)
        AIC: 479.2
        Number of Fisher Scoring iterations: 3
In [38]: # Variable CK, controlling for Daysrec:
        summary(fit_ck_days)
        glm(formula = Outcome ~ I(log(CK)) + Daysrec, family = binomial(),
            data = df
        Deviance Residuals:
           Min
                   10 Median
                                   30
                                          Max
         -2.195 -0.885 -0.559
                               1.039
                                        2.072
        Coefficients:
                    Estimate Std. Error z value Pr(>|z|)
         (Intercept)
                     4.0929
                                0.5887
                                         6.95 3.6e-12
                     -0.5939
                                0.0805
                                         -7.38 1.6e-13
         I(log(CK))
                     -0.1330
                                0.0643
        Daysrec
                                         -2.07
         (Dispersion parameter for binomial family taken to be 1)
            Null deviance: 546.64 on 409 degrees of freedom
        Residual deviance: 465.95 on 407 degrees of freedom
           (25 observations deleted due to missingness)
        AIC: 471.9
        Number of Fisher Scoring iterations: 3
In [25]: # Regress CK on Daysrec to see whether the time when the
        # blood measurement was taken matters.
         resp var pwr <- -0.2
```

```
Call:
         lm(formula = I(CK^resp_var_pwr) ~ Daysrec, data = df)
         Residuals:
                      1Q Median
            Min
                                      30
                                             Max
         -0.1441 -0.0554 -0.0129 0.0452 0.3475
         Coefficients:
                     Estimate Ctd Error + value Dr/al+1)
 In [ ]: | ### COMMENT:
         # Daysrec explains only 3.5% of the variation in CK.
In [35]: # Variable AST:
         fit_ast <- glm(Outcome ~ I(log(AST)), data= df, family= binomial())</pre>
         summary(fit_ast)
         Call:
         glm(formula = Outcome ~ I(log(AST)), family = binomial(), data = df)
         Deviance Residuals:
            Min
                    1Q Median
                                     30
                                            Max
         -1.889 -0.906 -0.535
                                1.020
                                          2.357
         Coefficients:
                     Estimate Std. Error z value Pr(>|z|)
                                   0.738
                                          7.33 2.4e-13
         (Intercept)
                       5.407
                      -1.089
                                   0.137
                                          -7.93 2.3e-15
         I(log(AST))
         (Dispersion parameter for binomial family taken to be 1)
             Null deviance: 569.75 on 428 degrees of freedom
         Residual deviance: 489.08 on 427 degrees of freedom
           (6 observations deleted due to missingness)
         AIC: 493.1
         Number of Fisher Scoring iterations: 4
In [39]: # Variable AST, controlling for Daysrec:
         fit_ast_days <- glm(Outcome ~ I(log(AST)) + Daysrec,</pre>
                             data= df, family= binomial())
         summary(fit_ast_days)
```

```
Call:
 In [ ]: ### COMMENTS:
         # We see from the above model that Daysrec and AST
         # are doing some of the same work in terms of
         # explaining the variance in the response variable.
         # The output suggests that we remove Daysrec
         # from the model. As already mentioned, I will keep
         # Daysrec in the final model to control for the
         # effects of this variable on the measurements of
         # both CK and AST.
         # Also, having a third dimension is likely to help
         # the k-means algorithm and give a boost to the tree
         # models examined further downstream.
In [35]: # Regress AST on Daysrec to see whether the time when the
         # blood measurement was taken matters.
         resp_var_pwr <- -0.2
         ast_days <- lm(I(AST^resp_var_pwr) ~ Daysrec, data= df)</pre>
         summary(ast_days)
         Call:
         lm(formula = I(AST^resp var pwr) ~ Daysrec, data = df)
         Residuals:
              Min
                        1Q Median
         -0.13600 -0.04345 -0.00063 0.04520 0.16391
         Coefficients:
                     Estimate Std. Error t value Pr(>|t|)
         (Intercept) 0.35054
                                0.00374 93.83 < 2e-16
                                 0.00119
                                           -6.04 3.4e-09
         Daysrec
                     -0.00719
         Residual standard error: 0.06 on 424 degrees of freedom
           (9 observations deleted due to missingness)
         Multiple R-squared: 0.0792, Adjusted R-squared: 0.077
         F-statistic: 36.4 on 1 and 424 DF, p-value: 3.43e-09
 In [ ]: ### COMMENT:
         # Daysrec explains around 8% of the variation in AST.
 In [8]: # See if Calving is needed when the other predictors are
         # in the model.
```

summary(fit_all)

data= df, family= binomial())

```
Call:
         glm(formula = Outcome \sim I(log(AST)) + I(log(CK)) + Daysrec +
             Calving, family = binomial(), data = df)
         Deviance Residuals:
                     1Q Median
         -2.087 -0.899 -0.516
                                0.975
                                          2.162
         Coefficients:
                     Estimate Std. Error z value Pr(>|z|)
         (Intercept)
                      4.9086 0.7918 6.20 5.7e-10
         I(log(AST))
                     -0.4898
                                  0.2208
                                           -2.22
                                                   0.0266
         I(log(CK))
                      -0.3798
                                  0.1233
                                           -3.08
                                                   0.0021
 In [ ]: ### COMMENTS:
         # The above model indicates that Calving is not going
         # to help us. This is OK because the overall aim of
         # this Part1 is not to find the best model for Outcome
         # given the original set of predictors. I just need
         # a subset of the predictors to work with; 3 is ideal,
         # because this greatly reduces the time needed to find
         # weights.
         # (It turns out that, for the purposes of showing the
         # value of a k-means hybrid model, I am actually better
         # off working with a set of predictors which make it
         # difficult for any classifier to have an accuracy score
         # greater than around 75-80%. As the accuracy score of
         # a classifier increases, the chances that the k-means
         # algorithm will be able to improve upon it decrease.)
In [10]: # Reduce our dataframe to include only the variables we
         # need. Then remove records with missing values.
         dfb <- df[, c("Outcome", "Daysrec", "CK", "AST")]</pre>
         dfb <- na.omit(dfb)</pre>
         nrow(dfb)
         408
```

Find best logistic regression model for predicting Outcome

```
Call:
glm(formula = Outcome ~ Daysrec + I(log(CK)) + I(log(AST)), family = binomial
data = dfb)

Deviance Residuals:

In []: ### COMMENTS:

# Each of Daysrec, CK, and AST are highly predictive of the
# outcome. AST and Daysrec do some of the same work in terms
# of predicting Outcome. Even so, it is important to control
# for this variable since CK and AST are blood measurements
# observed at different times for each cow. In this case,
# time is measured by Daysrec. We saw above that Daysrec
# explains around 3.5% of the variation in CK and around 8% of
# the variation in AST.

# It often helps to model with time squared. However,
# adding a quadratic, or the quadratic and cubic of
# Daysrec, to our model does not help.
```

f03 model diagnostics

Check f03 for overdispersion

```
In [63]: # If we have overdispersion, the standard errors for the
         # model coefficients will be too small.
         # We want the following value to NOT be considerably larger
         # than 1. If it is, then we have to use family= quasibinomial
         # in our modeling. (See p.311 of Kabacoff's "R in Action".)
         phi <- deviance(f03)/df.residual(f03)</pre>
         print(round(phi, 3))
         [1] 1.132
In [62]: # For a more accurate test, we need to also fit using family =
         # quasibinomial.
         f03_od <- glm(Outcome ~ Daysrec + I(log(CK)) + I(log(AST)),</pre>
                     data= dfb, family= quasibinomial)
In [64]: # See p.311 of Kabacoff's "R in Action". The null hypothesis
         \# is that the dispersion (phi) = 1. The alternative hypothesis
         # is that phi != 1. If the following p-value is small (e.g.,
         \# < 0.05), we have good evidence for rejecting the null
         # hypothesis.
         ans <- pchisq(summary(f03_od)$dispersion * f03$df.residual,</pre>
                        f03$df.residual, lower= FALSE)
         round(ans, 4)
         . - . . -
```

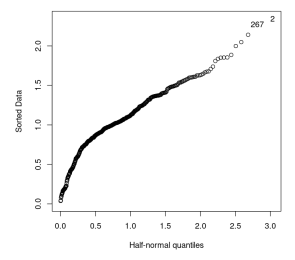
```
In [61]: # get_RsqrdDev is defined above. It computes the "proportion"
         # of deviance explained", an R-sqrd statistic for link= logit.
         f03_Rsqrd <- get_RsqrdDev(f03)</pre>
         print(f03 Rsqrd)
         # [1] 0.2588
         [1] 0.2588
In [43]: # Try different power transformations on the predictors to
         # see if we can boost the Rsqrd.
         f03 b <- glm(Outcome ~
                      I(Daysrec^1.5) +
                      I(CK^0.6) +
                      I(AST^{-0.33}),
                     data= dfb, family= binomial())
         summary(f03_b)
         print(get_RsqrdDev(f03_b))
         # [1] 0.2821
         Call:
         glm(formula = Outcome \sim I(Daysrec^1.5) + I(CK^0.6) + I(AST^-0.33),
             family = binomial(), data = dfb)
         Deviance Residuals:
            Min
                     1Q Median
                                     30
                                            Max
         -1.836 -0.934 -0.419 0.985
                                          2.345
         Coefficients:
                        Estimate Std. Error z value Pr(>|z|)
         (Intercept)
                        -0.56651 0.81313
                                             -0.70
                                                       0.486
         I(Daysrec^1.5) -0.04411
                                    0.02874
                                              -1.53
                                                       0.125
         I(CK^0.6)
                        -0.00851
                                    0.00214
                                              -3.97 7.1e-05
         I(AST^-0.33)
                         6.65170
                                    3.49487
                                               1.90
                                                       0.057
         (Dispersion parameter for binomial family taken to be 1)
             Null deviance: 543.76 on 407 degrees of freedom
         Residual deviance: 448.76 on 404 degrees of freedom
         AIC: 456.8
         Number of Fisher Scoring iterations: 5
         [1] 0.2821
```

406

```
In [44]: # Plot the residuals. Julian Faraway's half-normal plot can
    # be used to check for outliers. See p.46 of Faraway's
    # "Extending the Linear Model with R". The residuals plotted
    # here are the deviance residuals.

options(repr.plot.width= 6, repr.plot.height= 6)

# Function halfnorm is from package faraway.
halfnorm(residuals(f03_b))
```



```
In [45]: # Remove the 2 identified points. Faraway's halfnorm
# function identifies the points by their order in the
# dataframe and not by the rowname of the observation!

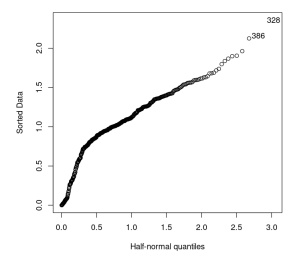
tmpdat <- dfb
nrow(tmpdat)
tmpdat <- tmpdat[c(-2, -267),]
nrow(tmpdat)</pre>
408
```

```
Call:
    glm(formula = Outcome ~ I(Daysrec^1.3) + I(CK^1) + I(AST^-0.3),
        family = binomial(), data = tmpdat)

Deviance Residuals:
        Min     10 Median     30 Max
        -1.798 -0.954 -0.315     0.979     2.360

Coefficients:
    [1] 0.3055

In [54]: options(repr.plot.width= 6, repr.plot.height= 6)
halfnorm(residuals(f03_b2))
```



```
In [55]: # Again, remove the 2 identified points. There are
# now 404 observations remaining.

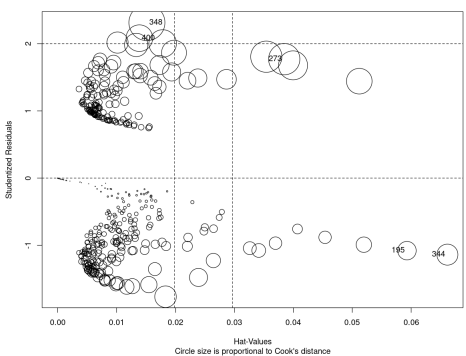
tmpdat <- tmpdat[c(-328, -386),]
nrow(tmpdat)</pre>
```

404

```
Call:
         glm(formula = Outcome \sim I(Daysrec^1.3) + I(CK^1) + I(AST^-0.3),
             family = binomial(), data = tmpdat)
         Deviance Residuals:
                     10 Median
                                            Max
         -1.817 -0.945 -0.264
                                0.968
                                          2.243
         Coefficients:
                         Estimate Std. Error z value Pr(>|z|)
         [1] 0.3251
In [68]: # Tweak the power transformations.
         f03_b4 <- glm(Outcome ~
                      I(Daysrec^1.5) +
                      I(CK^1.25) +
                      I(AST^-0.1),
                     data= tmpdat, family= binomial())
         summary(f03 b4)
         print(get_RsqrdDev(f03_b4))
         glm(formula = Outcome \sim I(Daysrec^1.5) + I(CK^1.25) + I(AST^-0.1),
             family = binomial(), data = tmpdat)
         Deviance Residuals:
            Min
                    1Q Median
                                     30
                                            Max
         -1.746 -0.959 -0.233 0.970 2.274
         Coefficients:
                         Estimate Std. Error z value Pr(>|z|)
                                    2.09e+00
                                              -2.46 0.01392
         (Intercept)
                        -5.13e+00
         I(Daysrec^1.5) -4.16e-02
                                    2.93e-02
                                               -1.42 0.15546
         I(CK^1.25)
                        -2.41e-05
                                    6.91e-06
                                              -3.49 0.00049
         I(AST^-0.1)
                                                2.71 0.00671
                         9.10e+00
                                    3.35e+00
         (Dispersion parameter for binomial family taken to be 1)
             Null deviance: 536.05 on 403 degrees of freedom
         Residual deviance: 425.55 on 400 degrees of freedom
         AIC: 433.6
         Number of Fisher Scoring iterations: 7
         [1] 0.3257
 In [ ]: ### COMMENT:
         # We started out with an Rsqrd of about 26%. With the
         # power transformations and the removal of 4 outliers,
         # we now have a model with an Rsqrd close to 32.6%.
In [601: ontions/repr plot width- 10 repr plot beight- 0)
```

	StudRes	Hat	CookD
	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
195	-1.0799	0.059290	0.012557
273	1.7705	0.038456	0.036109
344	-1.1393	0.066158	0.016244

Influence Plot for f03_b4



```
In [70]: # Remove obs. 348 from tmpdat. Here the point is identified
# by its rowname.

tmpdat02 <- tmpdat[which(!(rownames(tmpdat) %in% c(348))),]</pre>
```

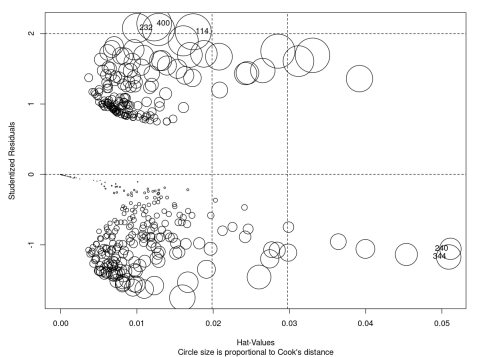
```
Call:
         glm(formula = Outcome \sim I(Daysrec^{1.5}) + I(CK^{1.2}) + I(AST^{-0.1}),
             family = binomial, data = tmpdat02)
         Deviance Residuals:
                     10 Madian
                                     30
            Min
         [1] 0.3352
In [74]: # Adjust the power transformations.
         f03_b6 <- glm(Outcome ~
                      Daysrec +
                      I(CK^1.2) +
                      I(AST^{-0.1}),
                     data= tmpdat02, family= binomial)
         summary(f03 b6)
         print(get_RsqrdDev(f03_b6))
         Call:
         glm(formula = Outcome \sim Daysrec + I(CK^1.2) + I(AST^-0.1), family = binomial
             data = tmpdat02)
         Deviance Residuals:
                  10 Median
                                     30
                                            Max
         -1.752 -0.960 -0.208 0.959
                                          2.139
         Coefficients:
                      Estimate Std. Error z value Pr(>|z|)
         (Intercept) -4.60e+00 2.16e+00 -2.13 0.03306
         Daysrec
                    -1.06e-01
                               7.10e-02 -1.50 0.13403
         I(CK^1.2)
                    -4.25e-05
                                1.16e-05 -3.66 0.00025
         I(AST^-0.1) 8.37e+00
                               3.45e+00
                                           2.43 0.01519
         (Dispersion parameter for binomial family taken to be 1)
             Null deviance: 534.11 on 402 degrees of freedom
         Residual deviance: 420.14 on 399 degrees of freedom
         AIC: 428.1
         Number of Fisher Scoring iterations: 7
         [1] 0.3355
In [75]: nrow(tmpdat02)
         403
In [76]: # Update our working dataframe.
         dfb <- tmpdat02
In [77]: # Save the file.
```

```
data= dfb, family= binomial())
         summary(f03_b6)
         print(get_RsqrdDev(f03_b6))
         # [1] 0.3355
         glm(formula = Outcome \sim Daysrec + I(CK^1.2) + I(AST^-0.1), family = binomial
             data = dfb
         Deviance Residuals:
                                     30
                    1Q Median
                                            Max
         -1.752 -0.960 -0.208 0.959
                                          2.139
         Coefficients:
                      Estimate Std. Error z value Pr(>|z|)
         (Intercept) -4.60e+00
                                 2.16e+00
                                            -2.13 0.03306
         Daysrec
                     -1.06e-01
                                 7.10e-02
                                            -1.50 0.13403
         I(CK^1.2)
                                 1.16e-05
                     -4.25e-05
                                            -3.66 0.00025
         I(AST^-0.1) 8.37e+00
                                3.45e+00
                                             2.43 0.01519
         (Dispersion parameter for binomial family taken to be 1)
             Null deviance: 534.11 on 402 degrees of freedom
         Residual deviance: 420.14 on 399 degrees of freedom
         AIC: 428.1
         Number of Fisher Scoring iterations: 7
         [1] 0.3355
In [79]: # If we do not lower the power transformation on CK, the
         # anova on the model generates a warning message
         # (probability of 0 or 1 in glm.fit). I find later
         # that even this is not enough to prevent the anova
         # function from issuing a warning. I am not sure what
         # the source of the problem is, especially given that
         # no such warning is generated by glm itself. (It may
         # be that the anova function uses different default
         # settings for glm.fit.)
         f03_b7 <- glm(Outcome ~
                      Daysrec +
                      CK +
                      I(log(AST)),
                     data= dfb, family= binomial())
         summary(f03_b7)
         print(get_RsqrdDev(f03_b7))
         # [1] 0.3345
```

A data.frame: 5×3

		StudRes	Hat	CookD
		<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
1	14	2.0255	0.017385	0.028599
23	32	2.0792	0.010017	0.018825
2	40	-1.0569	0.051118	0.010145
34	44	-1.1725	0.050894	0.013265
40	00	2.1425	0.012260	0.026451

Influence Plot for f03_b7



```
In [82]: options(repr.plot.width= 6, repr.plot.height= 6)
halfnorm(residuals(f03_b7), labs=rownames(dfb))
```

```
In [83]: # If we remove the last 3 influential points that are also
# outliers, our dataset will have been trimmed from 408
# records down to 400, a reduction of nearly 2%.
# Let's see what kind of difference this makes for our
# linear model.

tmpdat <- dfb[which(!(rownames(dfb) %in% c(400, 232, 114))),]
nrow(tmpdat)</pre>
400
```

Call:

```
glm(formula = Outcome \sim Daysrec + CK + I(log(AST)), family = binomial,
             data = tmpdat)
         [1] 0.3565
In [85]: # Update our working dataframe.
         dfb <- tmpdat
In [86]: # Save the file. dfb has 400 records in it.
         write.csv(dfb, file="/home/greg/Documents/stat/github repos/cows/downer.csv",
                  row.names= TRUE)
         Check model f03_b8 for overdispersion
In [87]: # If the following value is close to 1, the dispersion is
         # close to what our model assumes it to be. Thus, no
         # need to correct for overdispersion.
         phi <- deviance(f03_b8)/df.residual(f03_b8)</pre>
         print(round(phi, 3))
         [1] 1.028
In [89]: # See p.311 of Kabacoff's "R in Action". The null hypothesis
         # is that the dispersion (phi) = 1. The alternative hypothesis
         # is that phi != 1. If the following p-value is small (e.g.,
         \# < 0.05), we have good evidence for rejecting the null
         # hypothesis.
         f03b8_od <- glm(Outcome ~ Daysrec + CK + I(log(AST)),
```

0.9884

round(ans, 4) # 0.9884 (p-value)

Using anova output, compute the importance of the f03 predictors

data= dfb, family= quasibinomial)

Here the importances of the predictors in the f03_b8 model are determined by their anova contributions. terms in the model and the order of the terms matters, I need to use 6 different models and average over anova contribution for each model.

This section is included because I am curious to see whether the weights we arrive at for our base k-mea only AST, CK, and Daysrec as predictors) are at all like these anova weights.

* * * * *

```
Analysis of Deviance Table
          Model: binomial, link: logit
          Response: Outcome
          Terms added sequentially (first to last)
                        Df Deviance Resid. Df Resid. Dev
          NULL
                                           399
In [91]: names(f03_anova)
           'Df' · 'Deviance' · 'Resid. Df' · 'Resid. Dev'
In [136]: rownames(f03_anova)
           'NULL' · 'Daysrec' · 'CK' · 'I(log(AST))'
In [92]: ss01 <- f03_anova$'Deviance'</pre>
          names(ss01) <- c("null", "Daysrec", "CK", "AST")</pre>
          ss01tot <- sum(as.numeric(ss01[2:4]))</pre>
          print(round(ss01tot, 4))
           [1] 121.17
In [93]: | dayrec_percent <- round(ss01["Daysrec"]/ss01tot, 4)</pre>
          ck_percent <- round(ss01["CK"]/ss01tot, 4)</pre>
          ast_percent <- round(ss01["AST"]/ss01tot, 4)</pre>
          ss01_contribs <- c(dayrec_percent, ck_percent, ast_percent)</pre>
          print(ss01_contribs)
          Daysrec
                         CK
                                AST
            0.1152 0.8474 0.0374
In [161]: # Function for gauging the importance of the predictors
          # in a model based on each predictor's anova contribution.
          # As currently written, this function will not work if our
          # model contains interaction terms. This function is
          # specific to the f03 series of models.
          get_f03contribs <- function(dat) {</pre>
               # NOTE: the columns in dat must have the power
               # transformations used in the model. The first
               # column of dat must be the model response variable.
               preds <- colnames(dat)[-1]</pre>
               resp <- colnames(dat)[1]</pre>
               # get all permutations of the predictors;
               n <- length(preds)</pre>
               perm_matrix <- gtools::permutations(n, n, preds)</pre>
```

```
# The anova function calls glm.fit in a way that it sometimes
                   # produces warnings: "glm.fit: fitted probabilities numerically 0 or
                   modl_anova <- suppressWarnings(anova(modl))</pre>
                   ss <- modl_anova$'Deviance'</pre>
                   names(ss) <- rownames(modl anova)</pre>
                   ss_total <- round(sum(as.numeric(ss[-1])), 2)</pre>
                   percents <- round(ss[-1]/ss_total, 6)</pre>
                   names(percents) <- names(ss)[-1]</pre>
                   # accumulate results
                   for(name in names(result)) {
                       result[name] <- as.numeric(result[name]) + as.numeric(percents[name])
                   }
               } # end of for-loop
               return(round(result/nrow(perm_matrix), 4))
In [139]: # Create the dataframe for use with get_f03contribs.
          tmpdat <- dfb
          tmpdat$AST <- log(tmpdat$AST)</pre>
          dim(tmpdat)
          colnames(tmpdat)
           400 · 4
           'Outcome' · 'Daysrec' · 'CK' · 'AST'
In [162]: # The following weights are obtained using all 400 records.
          # These weights will be tested in Section 2 to see if they
          # can improve upon the basic k-means model, i.e., one without
          # weights.
          ans <- get_f03contribs(tmpdat)</pre>
          print(ans)
                           CK
          # Daysrec
             0.0528 0.5646 0.3827
          print(sum(ans))
          # [1] 1.0001
          Daysrec
                        CK
                                AST
            0.0528 0.5646 0.3827
           [1] 1.0001
```

Comments on anova weights

The anova weights are significantly different from the best weights for the k-means model found below.

```
In []:
```

After models are refined, model performance measures are obtained using cross-validation scores acros

For model comparison I introduce what I call a "Type2" score. This is a weighted average of the f-score (I accuracy (40%). The intent was to have a single score for model comparison. But I find in the end that fo dataset it is better to have, at minimum, the f-score, accuracy, and Type2 for model comparison. In Part3 where there are 3 classification levels to work with, I stick with just the accuracy score.

Basic Functions

```
In [3]: # This function is also found in Section 1.
         # Function for computing "proportion of deviance explained", an
         # R-sqrd statistic for link= logit. See p.41 of Julian Faraway's # "Extending the Linear Model with R" (2006: Chapman & Hall).
         # This statistic is due to N. Nagelkerke (1991; see Faraway's
         # bibliography).
         get RsqrdDev <- function(modl) {</pre>
              n obs <- length(modl$fitted.values)</pre>
              ans <- (1-exp((modl$deviance - modl$null.deviance)/n_obs))/(1-exp(-modl$r</pre>
              return(round(ans, 4))
In [4]: # Function to compute f-score for a 2x2 confusion matrix.
         get_fscore <- function(mat) {</pre>
             mat[is.na(mat)] <- 0</pre>
              FN <- as.numeric(mat[2,1])</pre>
              TP <- as.numeric(mat[2,2])</pre>
              FP <- as.numeric(mat[1,2])</pre>
              recall <- TP/(TP + FN)</pre>
              precision <- TP/(TP + FP)</pre>
              f_score <- 2* (recall*precision)/(recall + precision)</pre>
              return(round(f_score, 4))
In [5]: |# Function to output a confusion matrix and the f-score
         # for that matrix (if it is 2x2).
         get_confusion <- function(preds, df_actual) {</pre>
              # df_actual is a one-column dataframe;
              # preds is a named vector of predictions;
              # preds is of type factor; it is assumed there
              # are at least 2 factor levels
              levs <- levels(preds)</pre>
              n_levs <- length(levs)</pre>
              if(n_levs== 1) { levs <- c('0', '1') }</pre>
              n_levs <- max(n_levs, 2)</pre>
              actual <- as.vector(df_actual[, 1])</pre>
              names(actual) <- rownames(df_actual)</pre>
```

```
# which should be classed as such
    for(rowlev in levs) {
        actlev_names <- names(actual[actual == rowlev])</pre>
        # columns are for the predicted values:
        for(collev in levs) {
             predlev_names <- names(preds[preds == collev])</pre>
             if(length(predlev names > 0)) {
                 datout[rowlev, collev] <- sum(predlev_names %in% actlev_names</pre>
        }
        nonrow_cols <- levs[!(levs %in% rowlev)]</pre>
        datout[rowlev, "class.error"] <- round(sum(as.vector(datout[rowlev, r</pre>
                                                   sum(as.vector(datout[rowlev, l
    }
    result$matrix <- datout
    if(n levs == 2) {
        result[[2]] <- get_fscore(as.matrix(datout))</pre>
        result$f_score <- NA
    return(result)
}
```

```
In [6]: # Function for identifying which cluster each record
# belongs to.

getCluster <- function(x, centers) {

    # x is a row of a dataframe; its columns need
    # to be in the same order as centers (a matrix
    # constructed from kmeans)

    cl_dist <- apply(centers, 1, function(y) sqrt(sum((x-y)^2)))
    return(which.min(cl_dist)[1])
}</pre>
```

```
In [7]: # Function to generate combination of parameters for gridSearch;
       # each combination must add to a number ~1. Returns a dataframe,
       # each row of which is a valid combination.
       # I re-factored this ftn using R's expand.grid ftn. expand.grid
       # takes more time to run. This is probably due to type-checking.
       # It appears that we also run out of memory more quickly when
       # using expand.grid. Thus, I avoid expand.grid in what follows.
       generate_combs <- function(arglist, tol=0.0001) {</pre>
           # arglist is a named list; each name is a column
           \# name of the dataframe which goes to k-means
           # this next section is an alternative to expand.grid
           # if(FALSE) {
           n_args <- length(arglist)</pre>
           param_vlens <- rep(NA, n_args)</pre>
```

```
for(j in 1:n_args) {
                 vect <- arglist[[j]]</pre>
                 val <- rep(vect, rep(cprod, length(vect)))</pre>
                 datout[, j] <- rep(val, n_rows/length(val))</pre>
                 cprod <- cprod*length(vect)</pre>
             }
             # } ## end of 'if(FALSE)'
             # datout <- expand.grid(arglist, KEEP.OUT.ATTRS= FALSE)</pre>
             # colnames(datout) <- names(arglist)</pre>
             row sums <- round(rowSums(datout), 4)</pre>
             names(row_sums) <- rownames(datout)</pre>
             tol <- tol
             row sums <- row sums[which((as.numeric(row sums) <= (1 + tol)) & (as.num€
             datout <- datout[names(row_sums),]</pre>
             return(datout)
 In [8]: # Function to constrain range of data between 0 and 1.
         range01 <- function(x) {(x - min(x))/(max(x) - min(x))}
 In [9]: # Function to constrain range of data between min x and max x.
         # This function is used to transform validation data.
         range02 <- function(x, min_x, max_x) \{(x - min_x)/(max_x - min_x)\}
In [10]: # Function returning mapping between clusters and
         # Outcome levels. We choose the mapping that yields
         # the best accuracy score.
         c1 toLevel 1 <- function(dat) {</pre>
             # Returns TRUE if cluster 1 maps to Outcome= 1 (survivors)
             # dat is a dataframe with 2 columns, c("Outcome", "cluster");
             # nrow(dat) = number of predictions from the model;
             # dat$Outcome = traindat$Outcome (from the calling function)
             # We find the correct mapping between cluster number and
             # Outcome level by computing accuracy scores for the different
             # valid mappings. We choose the mapping with the best
             # accuracy score.
             tbl <- as.matrix(table(dat$Outcome, as.factor(dat$cluster)))
             # The colnames of tbl refer to the names of the clusters.
             # With only 2 levels for Outcome, we need only 2 scores.
             scores <- rep(NA, 2)</pre>
             # First possibility: cluster 1 maps to non-survivors
             tmpdat <- dat</pre>
             tmpdat[which(tmpdat$cluster== 1),]$Outcome <- 0</pre>
             tmpdat[which(tmpdat$cluster== 2),]$Outcome <- 1</pre>
```

```
preds <- as.factor(tmpdat$0utcome)
names(preds) <- rownames(tmpdat)
ans <- get_confusion(preds, dat[, "Outcome", drop=FALSE])
scores[2] <- ans[[2]]
return(scores[1] <= scores[2])
}</pre>
```

Optimization functions for random forest and gradient boostir models

```
In [11]: # Function for obtaining average of confusion matrix
          # f-score and percent correctly answered. This function
          # is called from get_cvScore and is used to find the best
          # parameters for the random forest and gradient boosting
          # models.
          get_Type2_rfgb <- function(traindat, testdat, classifier, ntrees,</pre>
                                        shrinkage) {
              if(classifier == 'randomforest') {
                   rfmod <- randomForest(I(as.factor(Outcome)) ~ .,</pre>
                                          data= traindat, ntree= ntrees,
                                          mtry= 1, nodesize= 1)
                  preds <- predict(rfmod, newdata= testdat, type="response")</pre>
                  ans <- get_confusion(preds, testdat[, "Outcome", drop=FALSE])</pre>
              if(classifier == 'gradientboost') {
                  gbmod <- suppressMessages(gbm(Outcome ~ ., data= traindat, n.trees= r</pre>
                                                   distribution= "bernoulli", shrinkage= s
                  preds <- suppressMessages(predict(gbmod, newdata= testdat, type="resr</pre>
                  preds transf <- preds</pre>
                  names(preds transf) <- rownames(testdat)</pre>
                  preds_transf[which(preds_transf >= 0.5)] <- 1</pre>
                  preds_transf[which(preds_transf < 0.5)] <- 0</pre>
                  preds_transf <- as.factor(preds_transf)</pre>
                  ans <- get_confusion(preds_transf, testdat[, "Outcome", drop=FALSE])</pre>
              }
              # Type2 score is a weighted average of accuracy and
              # the f-score.
              mat <- as.matrix(ans[[1]])</pre>
              percent_correct <- sum(diag(mat))/floor(sum(mat))</pre>
              result \leftarrow round((0.4 * percent_correct + 0.6 * ans[[2]]), 4)
              return(result)
          }
```

```
In [12]: # Function to obtain a cross-validation score, averaging the # Type2 scores of the folds. Valid values for the classifier # argument are: 'randomforest' and 'gradientboost'.
```

```
stopifnot(sum(segmentsv) == nrow(dat))
              # shuffle dat
              set.seed(seed)
              smp <- sample(rownames(dat), nrow(dat), replace= FALSE)</pre>
              dat <- dat[smp,]</pre>
              row_list <- vector("list", length=folds)</pre>
              names(row_list) <- as.character(1:folds)</pre>
              startpt <- 1
              for(i in 1:folds) {
                   endpt <- startpt + segmentsv[i] - 1</pre>
                   stopifnot(endpt <= nrow(dat))</pre>
                   row list[[i]] <- rownames(dat)[startpt:endpt]</pre>
                   startpt <- endpt + 1</pre>
              }
              train_list <- test_list <- vector("list", length= folds)</pre>
              for(j in 1:folds) {
                   valdat <- dat[row_list[[j]],]</pre>
                   traindat <- dat[which(!(rownames(dat) %in% rownames(valdat))),]</pre>
                   stopifnot((length(rownames(traindat)) + length(rownames(valdat))) ==
                   test_list[[j]] <- valdat</pre>
                   train_list[[j]] <- traindat</pre>
              }
              # With only 5 folds, we need only 5 cores.
              scores <- mcmapply(get_Type2_rfgb, train_list, test_list,</pre>
                                   MoreArgs= list(classifier= classifier,
                                                    ntrees= ntrees, shrinkage= shrinkage),
                                   SIMPLIFY=TRUE, mc.cores=5)
              # The average is of Type2 scores.
              return(round(mean(scores), 5))
In [13]: # Since the seed value has such a big effect on the results,
          # I take the average over a number of seeds.
          avg_seed_scores <- function(seed_vector, traindat, classifier,</pre>
                                         n_trees, shrinkage= 0.01, folds= 5) {
              seed_len <- length(seed_vector)</pre>
              outv <- rep(NA, seed_len)</pre>
              for(i in 1:seed_len) {
                   seed <- seed_vector[i]</pre>
```

In []:

```
# the cross-val scores.
datout <- rep(NA, 2 * tree_len * shrink_len)</pre>
dim(datout) <- c((tree_len * shrink_len), 2)</pre>
datout <- as.data.frame(datout)</pre>
colnames(datout) <- c("params", "Type2")</pre>
datout$params <- ""
index <- 0
for(i in 1:tree_len) {
    n_trees <- ntree_vector[i]</pre>
    if(classifier== 'gradientboost') {
         for(j in 1:shrink_len) {
             index <- index + 1
             shrinkage <- shrinkage_vector[j]</pre>
             param_string <- paste(as.character(n_trees),</pre>
                                     as.character(shrinkage), sep= "--")
             datout$params[index] <- param_string</pre>
             datout$Type2[index] <- avg_seed_scores(seed_vector, traindat,</pre>
                                                          folds=folds, shrinkac
        }
    }
    if(classifier== 'randomforest') {
        index <- index + 1</pre>
        datout$params[index] <- as.character(n_trees)</pre>
        datout$Type2[index] <- avg_seed_scores(seed_vector, traindat, cla</pre>
                                                     n_trees, folds= folds)
    }
}
return(datout)
```

Logistic regression: final model (g03)

When observations with missing values are removed from the original 435 record dataset, we are left witl Section 1 above I then constructed a linear model on these 408 records and found 8 of them to be outlied model), some of which were also influential. These 8 records were removed from the dataset. In this sect different models constructed from the 400 remaining records.

The removal of the model-based outliers may favor the logistic regression model over the other models logistic, there is often an overlap between variable-relative outliers and model-based outliers.

```
Call:
         glm(formula = Outcome \sim Daysrec + CK + I(log(AST)), family = binomial,
             data = dat)
         Deviance Residuals:
                    1Q Median
                                      30
         -1.752 -0.943 -0.186
                                0.943
                                           2.160
         Coefficients:
                      Estimate Std. Error z value Pr(>|z|)
         (Intercept) 2.89e+00 1.01e+00
                                            2.85 0.0044
         Daysrec
                     -1.03e-01
                                 7.19e-02
                                            -1.44
                                                    0.1507
                                            -4.02 5.9e-05
         CK
                     -3.15e-04
                                 7.84e-05
         I(log(AST)) -4.55e-01
                                2.15e-01
                                            -2.11 0.0346
         (Dispersion parameter for binomial family taken to be 1)
             Null deviance: 528.22 on 399 degrees of freedom
         [1] 0.3565
In [27]: preds <- predict(g03, newdata= dat, type="response")</pre>
         preds_transf <- preds</pre>
         preds_transf[which(preds_transf >= 0.5)] <- 1</pre>
         preds_transf[which(preds_transf < 0.5)] <- 0</pre>
         table(as.factor(preds_transf))
               1
         253 147
In [28]: # The following scores are not for comparing against the other
         # models. The value of the initial scores for each model is
         # to give us a general idea of what each model can do.
         preds transf <- as.factor(preds transf)</pre>
         ans <- get_confusion(preds_transf, dat[, "Outcome", drop=FALSE])</pre>
         print(ans$matrix)
         print(paste("f-score for model g03 (400) rcds): ", as.character(ans[[2]]), se
         # 0.6216
         # Accuracy = 72.0%
         # NOTE the even split between false positives and false negatives.
         # Type2 score is 0.6610
             0 1 class.error
         0 196 55
                    0.2191
         1 57 92
                       0.3826
         [1] "f-score for model g03 (400) rcds): 0.6216"
 In [ ]:
```

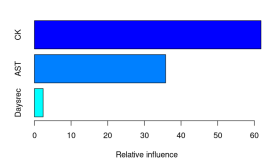
```
ans <- gridSearch02(seed_smp, dat, 'randomforest', tree_vector)</pre>
          (best_params <- ans[which(ans$Type2 == max(ans$Type2)),]$params)</pre>
          (best rf Type2 <- round(ans[which(ans$Type2 == max(ans$Type2)),]$Type2, 4))</pre>
          # 0.6194
          '160'
          0.6194
In [35]: # Refine the search. Test with 250 seeds.
          set.seed(7575)
          seed_smp <- sample(1:9999, 250, replace=FALSE)</pre>
          tree_vector <- c(160, 200, 240, 280)
          ans <- gridSearch02(seed_smp, dat, 'randomforest', tree_vector)</pre>
          (best_params <- ans[which(ans$Type2 == max(ans$Type2)),]$params)</pre>
          # '240'
          (best_rf_Type2 < - round(ans[which(ans$Type2 == max(ans$Type2)),]$Type2, 4))
          # 0.6210
          '240'
          0.621
In [36]: # Check the degree of variation in the Type2 scores.
          A data.frame: 4 x 2
           params
                   Type2
                   <dbl>
              160 0.61885
             200 0.61914
              240 0.62103
```

Best random forest classifier: rfclf_best

280 0.61994

Gradient boosting classifier

```
In [ ]: |# Here I use package gbm. When I work with the
         # wine dataset in Part 3, I use xgboost.
In [41]: # Run grid search to get parameters for the
         # gradient boosting model. This test is with 250 seeds.
         set.seed(7575)
         seed_smp <- sample(1:9999, 250, replace=FALSE)</pre>
         tree_vector <- c(80, 100, 120)
         shrinkage\_vector <- c(0.02, 0.03, 0.04)
         start <- Sys.time()</pre>
         ans <- gridSearch02(seed smp, dat, 'gradientboost', ntree vector=tree vector,
                              shrinkage_vector=shrinkage_vector, folds=5)
         stop <- Sys.time()</pre>
         round(stop - start, 2)
         # Time difference of 1.21 mins
          (best_params <- ans[which(ans$Type2 == max(ans$Type2, na.rm=TRUE)),]$params)</pre>
         # '100 - - 0.03'
         (best_gb_Type2 <- ans[which(ans$Type2 == max(ans$Type2, na.rm=TRUE)),]$Type2)</pre>
         # 0.6495
         Time difference of 1.21 mins
         '100--0.03'
         0.64955
In [42]: options(repr.plot.width= 6, repr.plot.height= 4)
         # The model is seed-dependent.
         set.seed(123)
         gbclf best <- gbm(Outcome ~ ., data= dat, n.trees= 100,
                            distribution= "bernoulli", shrinkage= 0.03)
          summarv(ahclf hest)
```



```
In [43]: # Initial model scores.
          preds <- suppressMessages(predict(gbclf_best, newdata= dat, type="response"))</pre>
          preds_transf <- preds</pre>
          names(preds_transf) <- rownames(dat)</pre>
          preds_transf[which(preds_transf >= 0.5)] <- 1</pre>
          preds_transf[which(preds_transf < 0.5)] <- 0</pre>
          preds_transf <- as.factor(preds_transf)</pre>
          ans <- get_confusion(preds_transf, dat[, "Outcome", drop=FALSE])</pre>
          print(ans$matrix)
          print(paste("f-score for gbclf best (400) rcds): ", as.character(ans[[2]]), s
          # f-score on training set: 0.6367
          # Accuracy on training set is 0.7375
          # Type2 score on training set is 0.6770
              0 1 class.error
          0 203 48
                         0.1912
            57 92
                         0.3826
          [1] "f-score for gbclf_best (400) rcds): 0.6367"
```

More stable scores for rfclf_best and gbclf_best

In order to get a better sense of how the random forest and gradient boosting models perform on the 400 to average the scores for these models over many different seeds.

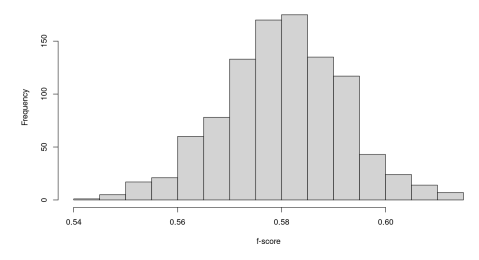
We see from the following results that gbclf_best performs much better than rfclf_best.

```
In [44]: # Get more stable scores for the best random forest model.

set.seed(1433)
seed_smp <- sample(1:9999, 1000, replace=FALSE)

datout <- rep(NA, 6 * length(seed_smp))
dim(datout) <- c(length(seed_smp), 6)</pre>
```

Distribution of f-scores for rfclf_best (400 rcds)

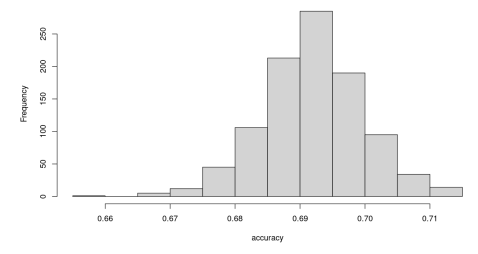


```
In [45]: # Summaries for rfclf_best.
    fn_avg <- mean(datout$FN)
    fp_avg <- mean(datout$FP)

    c(round(mean(datout$fscore), 4), round(mean(datout$Acc), 4),
        round(fn_avg, 2), round(fp_avg, 2))
# f-score: 0.5802
# accuracy: 0.6932
# false negatives: 65.43
# false positives: 55.49
# Average Type2 score is: 0.6254</pre>
```

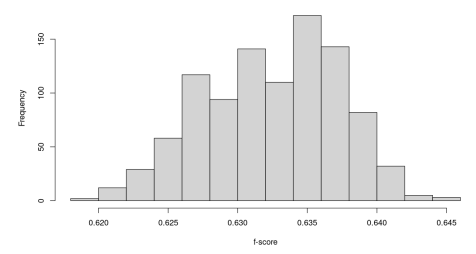
 $0.5802 \cdot 0.6932 \cdot 65.43 \cdot 55.49$

Distribution of accuracy scores for rfclf_best (400 rcds)

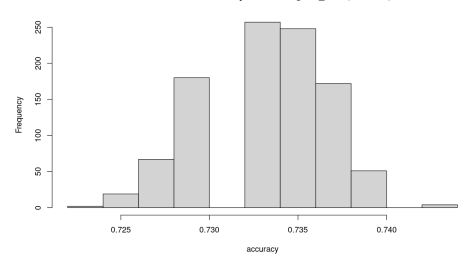


```
In [48]: # Get more stable scores for the best gradient
          # boosting model.
          set.seed(1433)
          seed smp <- sample(1:9999, 1000, replace=FALSE)</pre>
          datout <- rep(NA, 5 * length(seed_smp))</pre>
          dim(datout) <- c(length(seed_smp), 5)</pre>
          datout <- as.data.frame(datout)</pre>
          colnames(datout) <- c("seed", "fscore", "Acc", "FN", "FP")</pre>
          datout$seed <- seed smp</pre>
          for(i in 1:length(seed_smp)) {
              set.seed(seed_smp[i])
              gbmod <- gbm(Outcome ~ ., data= dat, n.trees= 100,</pre>
                              distribution= "bernoulli", shrinkage= 0.03)
              preds <- suppressMessages(predict(gbmod, newdata= dat, type="response"))</pre>
              preds_transf <- preds</pre>
              names(preds_transf) <- rownames(dat)</pre>
              preds_transf[which(preds_transf >= 0.5)] <- 1</pre>
              preds_transf[which(preds_transf < 0.5)] <- 0</pre>
              preds_transf <- as.factor(preds_transf)</pre>
              ans <- get_confusion(preds_transf, dat[, "Outcome", drop=FALSE])</pre>
              mat <- as.matrix(ans[[1]])</pre>
              percent_correct <- sum(diag(mat))/floor(sum(mat))</pre>
              datout[i, c("Acc")] <- round(percent_correct, 4)</pre>
              datout[i, c("fscore")] <- round(ans[[2]], 4)</pre>
              datout[i, c("FN")] <- as.numeric(mat[2,1])</pre>
              datout[i, c("FP")] <- as.numeric(mat[1,2])</pre>
          options(repr.plot.width= 10, repr.plot.height= 6)
          hist(datout$fscore, breaks=12, xlab="f-score"
                main="Distribution of f-scores for gbclf_best (400 rcds)")
```

Distribution of f-scores for gbclf_best (400 rcds)



Distribution of accuracy scores for gbclf_best (400 rcds)



SVM classifier

A support vector machine classifier using a radial basis function as the kernel.

```
In [35]: # Function to compute a Type2 score for an svm cv-fold.
          get Type2 svm <- function(traindat, valdat, gamma, cost) {</pre>
               # traindat and valdat need to be scaled
               train_scaled <- scale(traindat[, -1])</pre>
               train_centers <- attr(train_scaled, "scaled:center")
train_scales <- attr(train_scaled, "scaled:scale")</pre>
               train_scaled <- as.data.frame(cbind(traindat$0utcome, train_scaled),</pre>
                                                  row.names=rownames(traindat))
               colnames(train scaled) <- colnames(traindat)</pre>
               svmmod < - svm(I(as.factor(Outcome)) \sim ., data= train_scaled, gamma=gamma,
                                 cost=cost, scale=FALSE, kernel="radial")
               # Scale valdat.
               test_scaled <- scale(valdat[, -1], center=train_centers,</pre>
                                       scale=train_scales)
               test_scaled <- as.data.frame(cbind(valdat$Outcome,test_scaled),</pre>
                                                row.names=rownames(valdat))
               colnames(test_scaled) <- colnames(valdat)</pre>
               preds <- predict(svmmod, newdata= test scaled)</pre>
               ans <- table(preds, as.factor(valdat$0utcome))</pre>
               mat <- as.matrix(ans)</pre>
               mat[is.na(mat)] <- 0</pre>
               percent_correct <- sum(diag(mat))/floor(sum(mat))</pre>
               result <- round((0.4 * percent_correct + 0.6 * get_fscore(mat)), 4)
               return(result)
          }
```

```
In [36]: # This grid search searches for the best parameters for svm
          # modeling of the data.
          gridSearch svm <- function(seedv, dat, gammav, costv, folds=5) {</pre>
              gamma_len <- length(gammav)</pre>
              cost_len <- length(costv)</pre>
              # We need to capture the gridSearch parameters as well as
              # the cross-val scores.
              datout <- rep(NA, 2 * gamma_len * cost_len)</pre>
              dim(datout) <- c((gamma len * cost len), 2)</pre>
              datout <- as.data.frame(datout)</pre>
              colnames(datout) <- c("params", "Type2")</pre>
              datout$params <- ""
              # Divide dat by the number of folds to get a
              # size for each fold.
              segment_size <- round(nrow(dat)/folds)</pre>
              diff <- nrow(dat) - folds * segment_size</pre>
              last_seg_size <- segment_size + diff</pre>
              segmentsv <- c(rep(segment_size, (folds - 1)), last_seg_size)</pre>
              stopifnot(sum(segmentsv) == nrow(dat))
              index <- 0
              for(i in 1:gamma_len) {
                   gamma <- gammav[i]</pre>
```

```
seedv_len <- length(seedv)</pre>
             seed scores <- rep(NA, seedv len)
             for(h in 1:seedv_len) {
                 # shuffle dat
                 cur seed <- seedv[h]</pre>
                 set.seed(cur seed)
                 smp <- sample(rownames(dat), nrow(dat), replace= FALSE)</pre>
                 dat <- dat[smp,]</pre>
                 # Each element of row_list will be the rows we pick
                 \# out for one of the folds. E.g., the first element
                 # of row_list will contain the rows we want for the
                 # first fold, the second element of row list will
                 # contain the rows we want for the second fold, and
                 # so forth.
                 row list <- vector("list", length=folds)</pre>
                 names(row_list) <- as.character(1:folds)</pre>
                 startpt <- 1
                 for(k in 1:folds) {
                     endpt <- startpt + segmentsv[k] - 1</pre>
                     stopifnot(endpt <= nrow(dat))</pre>
                     row_list[[k]] <- rownames(dat)[startpt:endpt]</pre>
                     startpt <- endpt + 1</pre>
                 train_list <- test_list <- vector("list", length= folds)</pre>
                 for(k in 1:folds) {
                     testdat <- dat[row list[[k]],]</pre>
                     traindat <- dat[which(!(rownames(dat) %in% rownames(testc</pre>
                     stopifnot((length(rownames(traindat)) + length(rownames(t
                     test_list[[k]] <- testdat</pre>
                     train_list[[k]] <- traindat</pre>
                 }
                 # When there are only 5 folds, only 5 cores get used.
                 scores <- mcmapply(get Type2 svm, train list, test list,</pre>
                                      MoreArgs= list(gamma=gamma, cost=cost),
                                      SIMPLIFY= TRUE, mc.cores=5)
                 # For the current seed, store the average of the Type2
                 # scores, the average taken over the folds.
                 # We need na.rm=TRUE because one or more of the scores might
                 seed scores[h] <- round(mean(scores, na.rm=TRUE), 5)</pre>
             } ## end of for-loop, index h
             # Here I am taking an average of average scores. This
             # could be improved by simply taking a single average.
             datout$Type2[index] <- round(mean(seed scores), 5)</pre>
        } ## end of for-loop, index j
    } ## end of for-loop, index i
    return(datout)
}
```

```
In [20]: # Run grid search to get better parameters for the
# svm classifier.
```

```
round(stop - start, 2)
          # Time difference of 5.21 mins
          (best_params <- ans[which(ans$Type2 == max(ans$Type2)),]$params)</pre>
          # '0.01--20'
          (best Type2 <- ans[which(ans$Type2 == max(ans$Type2)),]$Type2)</pre>
          # 0.6573
          'Start time: 2021-07-21 10:55:25'
          Time difference of 5.21 mins
          '0.01--20'
          0.65729
In [39]: # Refine the search.
          set.seed(7543)
          seed vector <- sample(1:9999, 200, replace=FALSE)</pre>
          gamma_v \leftarrow seq(0.004, 0.01, by=0.002)
          cost_v < - seq(5, 20, by=5)
          start <- Sys.time()</pre>
          paste("Start time: ", start, sep="")
          ans <- gridSearch_svm(seed_vector, dat, gamma_v, cost v)</pre>
          stop <- Sys.time()</pre>
          round(stop - start, 2)
          # Time difference of 1.69 mins
          (best_params <- ans[which(ans$Type2 == max(ans$Type2)),]$params)</pre>
          # '0.006 - - 20 '
          (best_Type2 <- ans[which(ans$Type2 == max(ans$Type2)),]$Type2)</pre>
          # 0.6606
          'Start time: 2021-07-21 20:34:51'
          Time difference of 1.69 mins
          '0.006--20'
          0.66058
In [41]: # Construct an svm with the identified parameters.
          # We need to get probability estimates from the output.
          # So we set probability=TRUE.
          svm02 <- svm(I(as.factor(Outcome)) ~ ., data=svm_scaled, kernel="radial",</pre>
                        gamma= 0.006, cost= 20, scale=FALSE, probability=TRUE)
          pred <- fitted(svm02)</pre>
          (ans <- table(pred, as.factor(svm scaled$Outcome)))</pre>
          print(paste("f-score for 'best' svm classifier (400 rcds): ",
                       as.character(get_fscore(as.matrix(ans))), sep=""))
          # f-score: 0.6502
          # Accuracy: 0.7175
```

```
In [ ]:
```

Comparative cross-val scores for g03, svm02, and gbclf_best

I do not compute these scores for rfclf_best because, of the four models, it has the worst performance.

A summary table of these scores is presented at the end of Section 4.

```
In [12]: # This function is called by get_cvInfo. It returns a vector
          # of scores: f-score, accuracy, Type2, false negatives, and
          # false positives, in the stated order. The scores are for
          # the svm02 model.
          get_cvScores_svm02 <- function(traindat, valdat) {</pre>
               # traindat and valdat need to be scaled
               train_scaled <- scale(traindat[, -1])</pre>
               train_centers <- attr(train_scaled, "scaled:center")
train_scales <- attr(train_scaled, "scaled:scale")</pre>
               train_scaled <- as.data.frame(cbind(traindat$Outcome, train_scaled),</pre>
                                                 row.names=rownames(traindat))
               colnames(train_scaled) <- colnames(traindat)</pre>
               svmmod <- svm(I(as.factor(Outcome)) ~ ., data=train_scaled, kernel="radia"</pre>
                               gamma= 0.006, cost= 20, scale=FALSE, probability=TRUE)
               # Scale valdat.
               test_scaled <- scale(valdat[, -1], center=train_centers,</pre>
                                       scale=train_scales)
               test_scaled <- as.data.frame(test_scaled, row.names=rownames(valdat))</pre>
               preds <- predict(svmmod, newdata= test_scaled)</pre>
               ans <- table(preds, as.factor(valdat$0utcome))</pre>
               mat <- as.matrix(ans)</pre>
               mat[is.na(mat)] <- 0</pre>
               fscore <- round(get_fscore(mat), 4)</pre>
               acc <- round(sum(diag(mat))/floor(sum(mat)), 4)</pre>
               type2 <- round((0.4 * acc + 0.6 * fscore), 4)
               FN <- as.numeric(mat[2,1])</pre>
               FP <- as.numeric(mat[1,2])</pre>
               return(c(fscore,acc,type2,FN,FP))
```

```
In [41]: # This function returns 5 cross-validation metrics for each
# seed in the seed vector, seedv.

# (This function needs to be upgraded so that it can accept a
# function as an argument and then call that function within
# mcmapply.)

get_cvInfo <- function(seedv, dat, folds=5) {
    seedv_len <- length(seedv)
    datout <- rep(NA, 5 * folds * seedv_len)</pre>
```

```
diff <- nrow(dat) - folds * segment_size</pre>
    last seg size <- segment size + diff</pre>
    segmentsv <- c(rep(segment_size, (folds - 1)), last_seg_size)</pre>
    stopifnot(sum(segmentsv) == nrow(dat))
    for(h in 1:seedv_len) {
        # shuffle dat
        cur seed <- seedv[h]</pre>
        set.seed(cur_seed)
        smp <- sample(rownames(dat), nrow(dat), replace= FALSE)</pre>
        dat <- dat[smp,]</pre>
        # Each element of row_list will be the rows we pick
        # out for one of the folds. E.g., the first element
        # of row_list will contain the rows we want for the
        # first fold, the second element of row_list will
        # contain the rows we want for the second fold, and
        # so forth.
        row list <- vector("list", length=folds)</pre>
        names(row_list) <- as.character(1:folds)</pre>
        startpt <- 1
        for(k in 1:folds) {
             endpt <- startpt + segmentsv[k] - 1</pre>
             stopifnot(endpt <= nrow(dat))</pre>
             row_list[[k]] <- rownames(dat)[startpt:endpt]</pre>
             startpt \leftarrow endpt + 1
        }
        train list <- test list <- vector("list", length= folds)
        for(j in 1:folds) {
             testdat <- dat[row_list[[j]],]</pre>
             traindat <- dat[which(!(rownames(dat) %in% rownames(testdat))),]</pre>
             stopifnot((length(rownames(traindat)) + length(rownames(testdat))
             test list[[j]] <- testdat</pre>
             train_list[[j]] <- traindat</pre>
        # When there are only 5 folds, only 5 cores get used.
        ### NOTE: the function on the right-hand side changes depending on
        ### the model for which we want summary info.
        scores <- mcmapply(get_cvScores_kmp1_g03_v02, train_list, test_list,</pre>
                             SIMPLIFY= "array", mc.cores=5)
        for(k in 1:folds) {
             row name <- paste(as.character(cur seed), k, sep="--")</pre>
             datout[row_name, 1:5] <- scores[, k]</pre>
    } ## end of for-loop, index h
    return(datout)
}
```

svm02 comparative cross-val scores

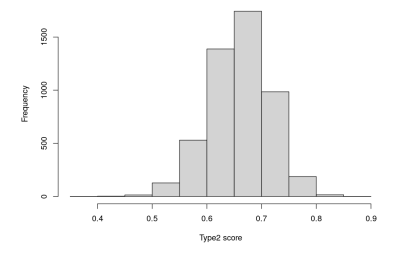
```
In [56]: # Get scores for the svm02 model.

set.seed(1913)
seed vector <- samnle(1.0000 1000 replace=FALSE)
```

```
In [57]:
          dim(dat_result)
          head(dat_result)
           5000 · 5
          A data.frame: 6 × 5
                                       FΝ
                                             FΡ
                  fscore
                          Acc Type2
                  <dbl>
                         <dbl>
                               <dbl> <dbl>
                                           <dbl>
           4782--1 0.5614 0.6875 0.6118
                                             16
           4782--2 0.6441 0.7375 0.6815
                                        16
                                              5
           4782--3 0.6667 0.6750 0.6700
                                        16
                                             10
           4782--4 0.6667 0.7125 0.6850
                                        15
                                              8
           4782--5 0.5806 0.6750 0.6184
                                        18
                                              8
           9275--1 0.6462 0.7125 0.6727
                                              9
In [58]: fscore_mean <- round(mean(dat_result$fscore, na.rm=TRUE), 4)</pre>
          fscore_sd <- round(sd(dat_result$fscore, na.rm=TRUE), 4)</pre>
          paste0("fscore mean: ", as.character(fscore_mean))
          # 0.6320
          paste0("fscore StdDev: ", as.character(fscore sd))
          # 0.0628
          summary(dat_result$fscore)
          'fscore mean: 0.632'
          'fscore StdDev: 0.0628'
             Min. 1st Qu. Median
                                        Mean 3rd Qu.
                                                          Max.
                                                                   NA's
            0.255
                     0.592
                                       0.632
                                               0.676
                                                         0.849
                              0.635
In [59]: Acc_mean <- round(mean(dat_result$Acc, na.rm=TRUE), 4)</pre>
          Acc_sd <- round(sd(dat_result$Acc, na.rm=TRUE), 4)</pre>
          paste0("accuracy mean: ", as.character(Acc_mean))
          paste0("accuracy StdDev: ", as.character(Acc_sd))
          summary(dat_result$Acc)
          'accuracy mean: 0.703'
          'accuracy StdDev: 0.0472'
             Min. 1st Qu. Median
                                        Mean 3rd Qu.
                                                          Max.
            0.438 0.675
                             0.700
                                       0.703 0.738
                                                         0.875
```

```
In [60]: Type2_mean <- round(mean(dat_result$Type2, na.rm=TRUE), 4)</pre>
         Type2_sd <- round(sd(dat_result$Type2, na.rm=TRUE), 4)</pre>
         paste0("Type2 mean: ", as.character(Type2_mean))
         paste0("Type2 StdDev: ", as.character(Type2_sd))
         # 0.0542
         summary(dat_result$Type2)
         'Type2 mean: 0.6604'
         'Type2 StdDev: 0.0542'
            Min. 1st Qu.
                           Median
                                      Mean 3rd Qu.
                                                       Max.
                                                               NA's
            0.378
                    0.626
                            0.662
                                     0.660
                                             0.698
                                                      0.859
In [61]: # Histogram of the Type2 scores for the svm02 model.
         options(repr.plot.width= 8, repr.plot.height= 6)
         hist(dat_result$Type2, breaks=10, xlab="Type2 score",
               main="Distribution of Type2 scores for the svm02 model")
```

Distribution of Type2 scores for the svm02 model



```
In [62]: FN_mean <- round(mean(dat_result$FN, na.rm=TRUE), 4)
    FN_sd <- round(sd(dat_result$FN, na.rm=TRUE), 4)
    paste0("FN mean: ", as.character(FN_mean))
# 14.53
    paste0("FN StdDev: ", as.character(FN_sd))
# 3.63
""
summary(dat_result$FN)</pre>
```

'FN mean: 14.5346'
'FN StdDev: 3.6265'

```
# 3.16
""
summary(dat_result$FP)

'FP mean: 9.229'
'FP StdDev: 3.1561'
"

Min. 1st Qu. Median Mean 3rd Qu. Max.
1.00 7.00 9.00 9.23 11.00 45.00

In []:
```

gbclf_best comparative cross-val scores

First change get_cvInfo so that it calls get_cvScores_gbclf.

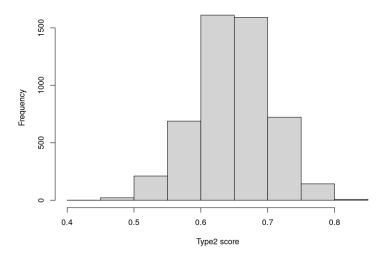
```
In [14]: # This function is called by get cvInfo. It returns a vector
          # of scores: f-score, accuracy, Type2, false negatives, and
          # false positives, in the stated order. The scores are for
          # the gbclf_best model.
          get_cvScores_gbclf <- function(traindat, valdat) {</pre>
              set.seed(123)
              gbmod <- gbm(Outcome ~ ., data= traindat, n.trees= 100,</pre>
                             distribution= "bernoulli", shrinkage= 0.03)
              preds <- suppressMessages(predict(gbmod, newdata= valdat, type="response"</pre>
              preds[which(preds >= 0.5)] <- 1
              preds[which(preds < 0.5)] <- 0
              names(preds) <- rownames(valdat)</pre>
              preds <- as.factor(preds)</pre>
              ans <- get_confusion(preds, valdat[, "Outcome", drop=FALSE])</pre>
              mat <- as.matrix(ans[[1]])</pre>
              mat[is.na(mat)] <- 0</pre>
              fscore <- round(ans[[2]], 4)
              acc <- round(sum(diag(mat))/floor(sum(mat)), 4)</pre>
              type2 <- round((0.4 * acc + 0.6 * fscore), 4)
              FN <- as.numeric(mat[2,1])</pre>
              FP <- as.numeric(mat[1,2])</pre>
              return(c(fscore,acc,type2,FN,FP))
In [66]: # Get summary scores for gbclf_best.
          set.seed(1913)
```

```
In [66]: # Get summary scores for gbclf_best.
set.seed(1913)
seed_vector <- sample(1:9999, 1000, replace=FALSE)
start <- Sys.time()
dat_result <- get_cvInfo(seed_vector, dat)
stop <- Sys.time()</pre>
```

A data.frame: 6 x 5

```
FΝ
                                             FP
                  fscore
                          Acc Type2
                  <dbl> <dbl>
                               <dbl> <dbl>
                                          <dbl>
           4782--1 0.4444 0.6250 0.5166
                                             10
           4782--2 0.6800 0.8000 0.7280
                                        7
                                              9
           4782--3 0.6377 0.6875 0.6576
                                             11
           4782--4 0.6769 0.7375 0.7011
                                             12
           4782--5 0.5763 0.6875 0.6208
                                             16
           9275--1 0.6316 0.7375 0.6740
                                              9
                                        12
In [68]: fscore_mean <- round(mean(dat_result$fscore, na.rm=TRUE), 4)</pre>
          fscore_sd <- round(sd(dat_result$fscore, na.rm=TRUE), 4)</pre>
          paste0("fscore mean: ", as.character(fscore_mean))
          # 0.6051
          paste0("fscore StdDev: ", as.character(fscore_sd))
          # 0.0665
          summary(dat_result$fscore)
          'fscore mean: 0.6051'
          'fscore StdDev: 0.0665'
             Min. 1st Qu. Median
                                        Mean 3rd Qu.
                                                         Max.
            0.353
                     0.561
                              0.607
                                       0.605
                                               0.654
                                                        0.814
In [69]: Acc_mean <- round(mean(dat_result$Acc, na.rm=TRUE), 4)</pre>
          Acc_sd <- round(sd(dat_result$Acc, na.rm=TRUE), 4)</pre>
          paste0("accuracy mean: ", as.character(Acc_mean))
          # 0.7135
          paste0("accuracy StdDev: ", as.character(Acc_sd))
          # 0.0448
          summary(dat_result$Acc)
          'accuracy mean: 0.7135'
          'accuracy StdDev: 0.0448'
             Min. 1st Qu. Median
                                        Mean 3rd Qu.
                                                         Max.
                                               0.750
            0.575
                     0.688
                              0.713
                                       0.714
                                                        0.863
In [70]: Type2_mean <- round(mean(dat_result$Type2, na.rm=TRUE), 4)</pre>
          Type2_sd <- round(sd(dat_result$Type2, na.rm=TRUE), 4)</pre>
          paste0("Type2 mean: ", as.character(Type2_mean))
          # 0.6485
          paste0("Type2 StdDev: ", as.character(Type2_sd))
          # 0.0552
          summary(dat result$Type2)
```

Distribution of Type2 scores for gbclf_best



```
In [72]: FN_mean <- round(mean(dat_result$FN, na.rm=TRUE), 4)
FN_sd <- round(sd(dat_result$FN, na.rm=TRUE), 4)
paste0("FN mean: ", as.character(FN_mean))
# 12.03
paste0("FN StdDev: ", as.character(FN_sd))
# 3.23
""
summary(dat_result$FN)</pre>
```

'FN mean: 12.0274'

'FN StdDev: 3.2325'

"

```
Min. 1st Qu. Median Mean 3rd Qu. Max. 2 10 12 12 14 26
```

```
In [73]: FP_mean <- round(mean(dat_result$FP, na.rm=TRUE), 4)
FP_sd <- round(sd(dat_result$FP, na.rm=TRUE), 4)
paste0("FP mean: ", as.character(FP_mean))
# 10.89
paste0("FP StdDev: ", as.character(FP_sd))
# 3.29
""
summary(dat_result$FP)</pre>
```

'FP mean: 10.8928'
'FP StdDev: 3.2897'

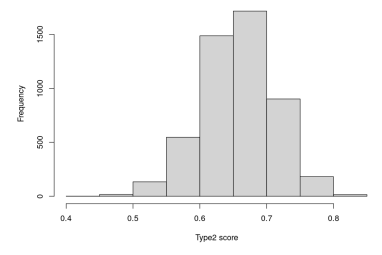
Again, first change get_cvInfo so that it calls get_cvScores_g03.

```
In [15]: # This function is called by get_cvInfo. It returns a vector
          # of scores: f-score, accuracy, Type2, false negatives, and
          # false positives, in the stated order.
          # (First alter get_cvInfo to call this function.)
          get_cvScores_g03 <- function(traindat, valdat) {</pre>
              g03mod <- suppressWarnings(glm(Outcome ~ Daysrec + CK + I(log(AST))),</pre>
                              data= traindat, family= binomial, singular.ok=TRUE,
                              epsilon= 1e-7, maxit=50))
              preds <- suppressWarnings(predict(g03mod, newdata= valdat, type="response"
              preds[which(preds >= 0.5)] <- 1
              preds[which(preds < 0.5)] <- 0
              names(preds) <- rownames(valdat)</pre>
              preds <- as.factor(preds)</pre>
              ans <- get_confusion(preds, valdat[, "Outcome", drop=FALSE])</pre>
              mat <- as.matrix(ans[[1]])</pre>
              mat[is.na(mat)] <- 0</pre>
              fscore <- round(ans[[2]], 4)
              acc <- round(sum(diag(mat))/floor(sum(mat)), 4)</pre>
              type2 <- round((0.4 * acc + 0.6 * fscore), 4)
              FN <- as.numeric(mat[2,1])</pre>
              FP <- as.numeric(mat[1,2])</pre>
              return(c(fscore,acc,type2,FN,FP))
In [76]: # Get summary scores for the g03 logistic model.
          set.seed(1913)
          seed_vector <- sample(1:9999, 1000, replace=FALSE)</pre>
          start <- Sys.time()</pre>
          dat_result <- get_cvInfo(seed_vector, dat)</pre>
          stop <- Sys.time()</pre>
          round(stop - start, 2)
          # Time difference of 31.94 secs
          Time difference of 31.94 secs
In [77]: dim(dat_result)
          head(dat_result)
          5000 · 5
          A data.frame: 6 × 5
                                             FP
                                       FΝ
                  fscore
                          Acc Type2
                               <dbl> <dbl>
                                          <dbl>
                  <dbl>
                        <dbl>
           4782--1 0.4615 0.6500 0.5369
                                       20
                                              8
           4782--2 0.7170 0.8125 0.7552
                                        5
                                             10
```

```
paste0("fscore mean: ", as.character(fscore_mean))
         paste0("fscore StdDev: ", as.character(fscore_sd))
         # 0.0646
         summary(dat_result$fscore)
         'fscore mean: 0.6183'
         'fscore StdDev: 0.0646'
            Min. 1st Qu. Median
                                     Mean 3rd Qu.
                                                      Max.
           0.353 0.576
                                     0.618 0.667
                                                     0.828
                            0.621
In [79]: Acc_mean <- round(mean(dat_result$Acc, na.rm=TRUE), 4)</pre>
         Acc_sd <- round(sd(dat_result$Acc, na.rm=TRUE), 4)</pre>
         paste0("accuracy mean: ", as.character(Acc_mean))
         # 0.7167
         paste0("accuracy StdDev: ", as.character(Acc_sd))
         # 0.0448
         summary(dat_result$Acc)
         'accuracy mean: 0.7167'
         'accuracy StdDev: 0.0448'
            Min. 1st Qu. Median
                                      Mean 3rd Qu.
                                                      Max.
            0.550 0.688
                           0.713
                                     0.717 0.750
                                                     0.875
In [80]: Type2_mean <- round(mean(dat_result$Type2, na.rm=TRUE), 4)</pre>
         Type2_sd <- round(sd(dat_result$Type2, na.rm=TRUE), 4)</pre>
         paste0("Type2 mean: ", as.character(Type2_mean))
         # 0.6577
         paste0("Type2 StdDev: ", as.character(Type2_sd))
         # 0.0543
         summary(dat_result$Type2)
         'Type2 mean: 0.6577'
         'Type2 StdDev: 0.0543'
            Min. 1st Qu. Median
                                      Mean 3rd Qu.
                                                      Max.
            0.447 0.621 0.659
                                     0.658 0.695
                                                     0.847
```

```
In [81]: # Histogram of the Type2 scores for g03.
    options(repr.plot.width= 8, repr.plot.height= 6)
    hist(dat_result$Type2, breaks=10, xlab="Type2 score",
        main="Distribution of Type2 scores for the g03 logistic model")
```

Distribution of Type2 scores for the g03 logistic model



```
In [82]: FN_mean <- round(mean(dat_result$FN, na.rm=TRUE), 4)
FN_sd <- round(sd(dat_result$FN, na.rm=TRUE), 4)
paste0("FN mean: ", as.character(FN_mean))
# 11.24
paste0("FN StdDev: ", as.character(FN_sd))
# 3.04
""
summary(dat_result$FN)</pre>
```

'FN mean: 11.2392'

'FN StdDev: 3.038'

"

Min. 1st Qu. Median Mean 3rd Qu. Max. 2.0 9.0 11.0 11.2 13.0 23.0

```
In [83]: FP_mean <- round(mean(dat_result$FP, na.rm=TRUE), 4)
    FP_sd <- round(sd(dat_result$FP, na.rm=TRUE), 4)
    paste0("FP mean: ", as.character(FP_mean))
# 11.43
    paste0("FP StdDev: ", as.character(FP_sd))
# 3.34
""
summary(dat_result$FP)</pre>
```

'FP mean: 11.426'

'FP StdDev: 3.3402'

has the best accuracy score (0.7167), edging out gbclf_best (0.7135). A Type2 score is 60% of the f-scoraccuracy. svm02's Type2 is 0.6604; g03's Type2 is 0.6577; gbclf_best's Type2 is 0.6485.

On average, svm02 has the fewest false positives but the most false negatives. g03 tends to have an equipositives and false negatives.

A table of the above scores is found at the end of Section 4.

* * * * *

Section 3: A k-means base model

How does the k-means algorithm cluster the 400 observations for AST, CK, and Daysrec, and how well d clusters align with the levels of Outcome?

```
In [13]: # Transform data for plotting.

dat_plot <- dat
    dat_plot$AST <- log(dat_plot$AST)
    dat_plot$CK <- log(dat_plot$CK)</pre>
```

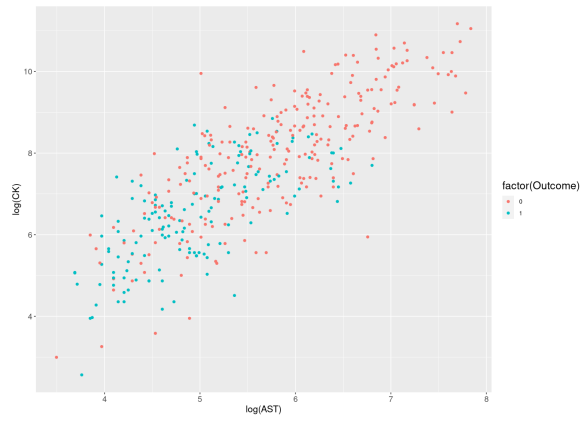
```
In [14]: # Plot data to see the degree of separation between
# records with Outcome = 1 and those with Outcome = 0,
# ignoring for the moment the Daysrec dimension.

options(repr.plot.width= 12, repr.plot.height= 9)

p <- ggplot(dat_plot, aes(AST, CK, color= factor(Outcome))) +

geom_point(alpha= 1.0) +
    xlab("log(AST)") + ylab("log(CK)") +
    ggtitle("Outcome differences in CK and AST") +
    theme(axis.text= element_text(size = 12)) +
    theme(axis.title= element_text(size= 14)) +
    theme(title= element_text(size= 16))
p</pre>
```

Outcome differences in CK and AST



```
In []: ### COMMENT:
    # We see in the above plot that there is a degree of
    # separation in Outcome levels based on the AST and
    # CK blood measurements. These are the 2 most important
    # dimensions we have for predicting Outcome.
In [57]: ### COMMENT:
```

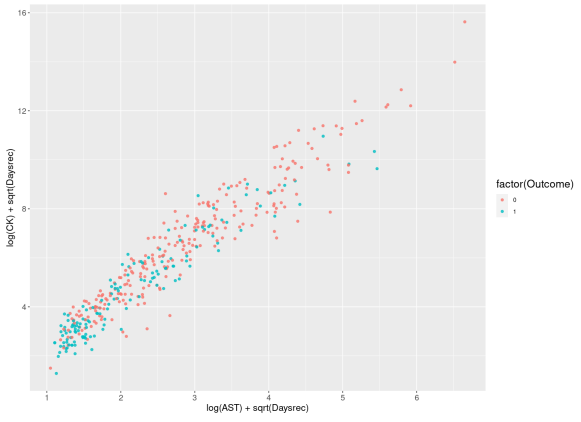
```
dat_plot$AST <- log(dat_plot$AST) * 0.3 * sqrt(1 + dat_plot$Daysrec)
dat_plot$CK <- log(dat_plot$CK) * 0.5 * sqrt(1 + dat_plot$Daysrec)

options(repr.plot.width= 12, repr.plot.height= 9)

p <- ggplot(dat_plot, aes(AST, CK, color= factor(Outcome))) +

geom_point(alpha= 0.8) +
    xlab("log(AST) + sqrt(Daysrec)") + ylab("log(CK) + sqrt(Daysrec)") +
    ggtitle("Outcome differences in CK and AST") +
    theme(axis.text= element_text(size = 12)) +
    theme(axis.title= element_text(size= 14)) +
    theme(title= element_text(size= 16))
p</pre>
```

Outcome differences in CK and AST



```
In [59]: # Try a different transformation.

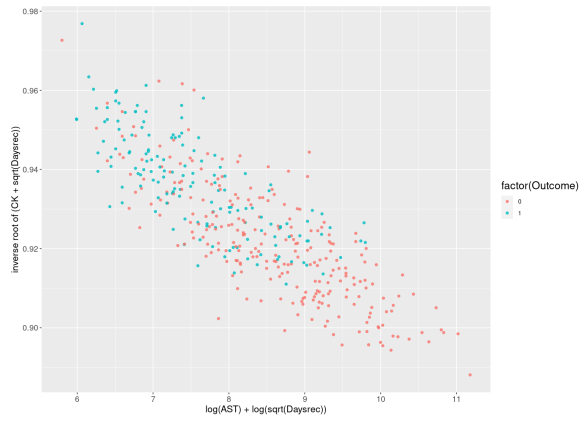
dat_plot <- dat
    dat_plot$AST <- log(dat_plot$AST * 10 * sqrt(1 + dat_plot$Daysrec))
    dat_plot$CK <- (dat_plot$CK * (0.8 * sqrt(1 + dat_plot$Daysrec)))^-0.01</pre>
```

```
In [60]: options(repr.plot.width= 12, repr.plot.height= 9)

p <- ggplot(dat_plot, aes(AST, CK, color= factor(Outcome))) +

geom_point(alpha= 0.8) +
    xlab("log(AST) + log(sqrt(Daysrec))") + ylab("inverse root of (CK + sqrt(Daysrec))") +
    ggtitle("Outcome differences in CK and AST") +
    theme(axis.text= element_text(size= 12)) +
    theme(axis.title= element_text(size= 14)) +
    theme(title= element_text(size= 16))
p</pre>
```

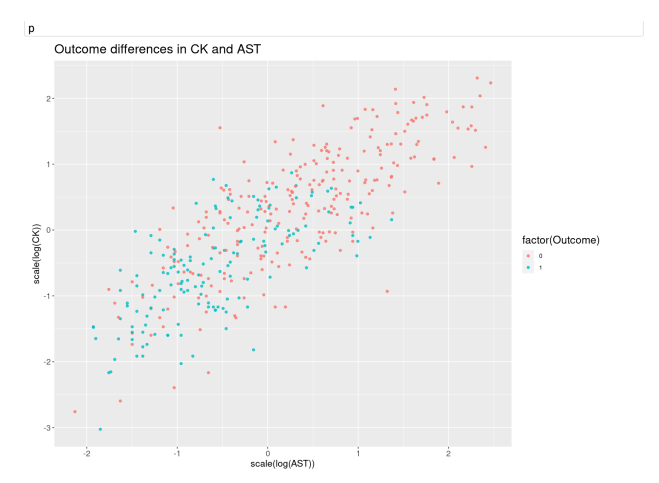
Outcome differences in CK and AST



```
In [20]: # Transform the data as it will be transformed
# for k-means. HTF in "The Elements of Statistical Learning"
# point out that there are times when scaling makes it MORE
# difficult for clustering algorithms to separate the data
# as we would like, or expect. See Figure 14.5 on p.506.

dat_plot <- dat
dat_plot$AST <- log(dat_plot$AST)
dat_plot$CK <- log(dat_plot$CK)

dat_plot$AST <- scale(dat_plot$AST)
dat_plot$CK <- scale(dat_plot$CK)</pre>
```



Initial k-means model

```
# Remove Outcome.
          set.seed(1233)
          fit_km <- kmeans(df_scaled[, -1], 2, iter.max = 50, nstart = 30)</pre>
          print(fit_km$size)
          # [1] 216 184
           [1] 183 217
In [20]: datout <- as.data.frame(cbind(df_scaled$Outcome, fit_km$cluster))
    colnames(datout) <- c("Outcome", "cluster")</pre>
          rownames(datout) <- rownames(df_scaled)</pre>
          head(datout)
          A data.frame: 6 \times 2
              Outcome cluster
                 <dbl>
                        <dbl>
           3
                    0
                           1
                           1
                           2
                    0
                           2
                    0
                           2
In [21]: # max_vals identifies the Outcome level we would expect
          # each cluster to map to.
           (mat <- as.matrix(table(datout$Outcome, as.factor(datout$cluster))))</pre>
          max_vals <- apply(mat, MARGIN=2, which.max); print(max_vals)</pre>
                 1
             0 78 173
             1 105 44
          2 1
In [22]: # Use function c1_toLevel_1 to get the correct
          # mapping between clusters and Outcome levels.
          c1_to_Outcome1 <- c1_toLevel_1(datout)</pre>
          paste("Map cluster 1 to Outcome level 1? : ", c1_to_Outcome1, sep="")
          'Map cluster 1 to Outcome level 1?: TRUE'
In [23]: tmpdat <- datout</pre>
          tmpdat[which(tmpdat$cluster== 1),]$Outcome <- 1</pre>
          tmpdat[which(tmpdat$cluster== 2),]$Outcome <- 0</pre>
          dim(tmpdat)
           400 O
```

```
In [25]: # Get initial set of scores for the k-means model.

preds <- as.factor(tmpdat$0utcome)
names(preds) <- rownames(tmpdat)
ans <- get_confusion(preds, df_scaled[, "Outcome", drop=FALSE])
print(ans$matrix)

""

print(paste("f-score for k-means base model (400 rcds): ", as.character(ans[]
# f-score: 0.6325

# Accuracy: 0.6950

# Type2: 0.6575</pre>

0 1 class.error
0 173 78 0 3108
```

Comments regarding scores for k-means base model

In the next section on weights, the base model (without weights) is run against 2500 folds; the average or is 0.6533. This score beats that of our best gradient boosting model, gbclf_best (which has a comparative score of 0.6485). The latter is supervised learning; the former is unsupervised.

The g03 model has a comparative Type2 cross-val score of 0.6577, only 0.44 percentage points higher the score. What is notable about g03 is that, among all of the models surveyed thus far, it has the highest account (0.7167, comparative cross-val) and the second-highest f-score.

In short, the unsupervised k-means model does very well as a classifier for this dataset. It is already com very best models. This is a good indicator that we can use k-means to construct a better classifier than a have already surveyed. We in fact see in the next section that we only have to add weights to the model (k-means is now being used in a supervised learning setting) to have a model with a higher Type2 score t svm02, or gbclf_best.

Find weights for k-means base model

Getting weights means running a grid search. Since we want our model to apply to unseen data, the weights established using cross-validation. Since the parameter space is fairly large, I evaluate the weights based "folds". In this example, I identify the best weights as those which yield the best average Type2 score. I u because it is based on accuracy, precision, and recall; it is thus a more general score than either accuracy f-score. For this dataset I have not yet tried to find weights based on accuracy score alone. Of course, if a criterion by which we wanted to judge our models, then the best weights should be identified as those whighest accuracy score.

Whereas the above k-means base model is unsupervised, the k-means base model with weights is const supervised learning.

* * * * *

```
# below. (Here the names are: AST, CK, and Daysrec.)
####################################
# Transform and scale training set data for the
# k-means model.
traindat$AST <- log(traindat$AST)</pre>
traindat$CK <- log(traindat$CK)</pre>
traindat$Daysrec <- sqrt(traindat$Daysrec)</pre>
traindat_scaled <- scale(traindat[, -1], center=TRUE, scale=TRUE)</pre>
centers <- attr(traindat_scaled, "scaled:center")
scales <- attr(traindat_scaled, "scaled:scale")</pre>
###############################
# Apply weights to traindat. The sqrt should have
# been taken in the calling function.
cols <- names(wghts)</pre>
df2 <- t(t(traindat_scaled[, cols]) * as.numeric(wghts[cols]))</pre>
traindat_wghts <- as.data.frame(df2, row.names=rownames(traindat))</pre>
colnames(traindat_wghts) <- cols</pre>
###############################
# Prepare valdat.
# Transform and scale valdat.
valdat$AST <- log(valdat$AST)</pre>
valdat$CK <- log(valdat$CK)</pre>
valdat$Daysrec <- sqrt(valdat$Daysrec)</pre>
valdat_scaled <- scale(valdat[, -1], center=centers, scale=scales)</pre>
valdat_scaled <- as.data.frame(valdat_scaled, row.names=rownames(valdat))</pre>
# Apply weights to valdat. (We want valdat to look exactly like
# traindat. The weights act as a transformation of the data.)
df2 <- t(t(valdat_scaled[, cols]) * as.numeric(wghts[cols]))</pre>
valdat wghts <- as.data.frame(df2, row.names=rownames(valdat))</pre>
colnames(valdat_wghts) <- cols</pre>
##################################
# Construct k-means model.
kmod <- suppressWarnings(kmeans(traindat wghts, 2, iter.max = 50, nstart=
# See how the clusters are associated with Outcome.
dfout <- as.data.frame(cbind(traindat$Outcome, kmod$cluster),</pre>
                         row.names=rownames(traindat))
colnames(dfout) <- c("Outcome", "cluster")</pre>
c1 to Outcome1 <- c1 toLevel 1(dfout)</pre>
###############################
# Apply the k-means model to valdat_wghts.
# Each element of the following list is a row of valdat_wghts.
valdat_asList <- split(valdat_wghts[, colnames(kmod$centers)],</pre>
                         seq(nrow(valdat_wghts)))
stm list - ...stam/#list# lamath_ mma./..sldat\\
```

```
valdat_wghts$cluster <- as.numeric(cluster_assgns)</pre>
              valdat_wghts$pred_Outcome <- NA</pre>
              if(c1_to_Outcome1) {
                  valdat_wghts[which(valdat_wghts$cluster==1),]$pred_Outcome <- 1</pre>
                  valdat_wghts[which(valdat_wghts$cluster==2),]$pred_Outcome <- 0</pre>
                  valdat_wghts[which(valdat_wghts$cluster==1),]$pred_Outcome <- 0</pre>
                  valdat_wghts[which(valdat_wghts$cluster==2),]$pred_Outcome <- 1</pre>
              }
              # Generate confusion matrix for the k-means clusters and
              # the corresponding f-score.
              preds <- as.factor(valdat wghts$pred Outcome)</pre>
              names(preds) <- rownames(valdat)</pre>
              ans <- get_confusion(preds, valdat[, "Outcome", drop=FALSE])</pre>
              # The result returned is a Type2 score (which is a mixture
              # of accuracy and f-score).
              mat <- as.matrix(ans[[1]])</pre>
              mat[is.na(mat)] <- 0</pre>
              percent correct <- sum(diag(mat))/floor(sum(mat))</pre>
              result <- round((0.4 * percent\_correct + 0.6 * ans[[2]]), 6)
              return(result)
In [13]: # This grid search searches for the best set of weights to use
         # in our k-means clustering model. The best weights are those
         # which generalize best to the validation set. So we look for
         # the best cross-validation score.
         # Because our training set is so small---only 400 records---we
         # need to run the gridSearch over many seeds. Otherwise, we
         # will not get a meaningful result.
         gridSearch06 <- function(seed_vector, dat, df_params, folds=5) {</pre>
              datout <- rep(NA, 2*nrow(df_params))</pre>
              dim(datout) <- c(nrow(df_params), 2)</pre>
              datout <- as.data.frame(datout)</pre>
              colnames(datout) <- c("row", "Type2")</pre>
              datout$row <- rownames(df_params)</pre>
              # We want the sqrt of the weights.
              df_params <- df_params^0.5</pre>
              params_rows <- rownames(df_params)</pre>
              ###############################
              # Partition the data into folds.
              # divide dat by the number of folds
              segment_size <- round(nrow(dat)/folds)</pre>
              diff <- nrow(dat) - folds * segment_size</pre>
              last_seg_size <- segment_size + diff</pre>
              segmentsv <- c(rep(segment size, (folds - 1)), last seg size)</pre>
              stopifnot(sum(segmentsv) == nrow(dat))
              # Create a dataframe, each row for a distinct seed.
              # Each column of the dataframe is for a distinct set
              # of weights. The entries in the cells are Type2
```

```
set.seed(cur_seed)
        smp <- sample(rownames(dat), nrow(dat), replace= FALSE)</pre>
        dat <- dat[smp,]</pre>
        # Each element of row_list will be the rows we pick
        # out for one of the folds. E.g., the first element
        # of row list will contain the rows we want for the
        # first fold, the second element of row_list will
        # contain the rows we want for the second fold, and
        # so forth.
        row_list <- vector("list", length=folds)</pre>
        names(row_list) <- as.character(1:folds)</pre>
        startpt <- 1
        for(i in 1:folds) {
             endpt <- startpt + segmentsv[i] - 1</pre>
             # stopifnot(endpt <= nrow(dat))</pre>
             row list[[i]] <- rownames(dat)[startpt:endpt]</pre>
             startpt <- endpt + 1</pre>
        }
        for(i in 1:nrow(df_params)) {
             cur_row <- params_rows[i]</pre>
             wghts <- as.numeric(df_params[i,])</pre>
             names(wghts) <- colnames(df_params)</pre>
             train_list <- test_list <- vector("list", length= folds)</pre>
             for(j in 1:folds) {
                 testdat <- dat[row list[[j]],]</pre>
                 traindat <- dat[which(!(rownames(dat) %in% rownames(testdat))</pre>
                 # stopifnot((length(rownames(traindat)) + length(rownames(tes
                 test_list[[j]] <- testdat</pre>
                 train_list[[j]] <- traindat</pre>
             # When there are only 5 folds, only 5 cores get used.
             ### NOTE: I change the following function call depending on
             ### the model I am scoring.
             scores <- mcmapply(get_cvScore_kmp1_g03, train_list, test_list,</pre>
                                 MoreArgs= list(wghts=wghts),
                                 SIMPLIFY= TRUE, mc.cores=5)
             # For the current seed, store the average of the Type2
             # scores, the average taken over the folds.
             df scores[as.character(cur seed), cur row] <- round(mean(scores),</pre>
        } # end of for-loop, index i
    } ## end of for-loop, index h
    # Compute the average over the seeds of the Type2 scores
    # obtained for each set of parameters in df params.
    datout$Type2 <- round(apply(df scores, MARGIN=2, mean), 5)</pre>
    return(datout)
}
```

```
In [33]: # There are 3 parameter lists to work with.
          lst <- vector("list", length= 3)
names(lst) <- c("AST","CK","Daysrec")</pre>
          lst[[1]] \leftarrow lst[[2]] \leftarrow lst[[3]] \leftarrow seq(0.15, 0.55, by=0.01)
          start <- Sys.time()</pre>
          dfc01 <- generate_combs(lst, tol=0.0001)</pre>
          stop <- Sys.time()</pre>
          # round(stop - start, 2)
          dim(dfc01)
          # 1236
           1236 · 3
In [34]: # Test on a sample of 10.
          set.seed(42)
          smp <- sample(rownames(dfc01), 10, replace=FALSE)</pre>
          tst_params <- dfc01[smp,]</pre>
In [35]: # Find the best weights of those in tst_params.
          set.seed(1233)
          seed_vector <- sample(1:9999, 11, replace=FALSE)</pre>
          start <- Sys.time()</pre>
          dat_result <- gridSearch06(seed_vector, dat, tst_params)</pre>
          stop <- Sys.time()</pre>
          round(stop - start, 2)
          # Time difference of 18 secs (for 10 rows, 11 seeds)
          Time difference of 19.91 secs
In [36]: best_params <- dat_result[which(dat_result$Type2 ==</pre>
                                                max(dat result$Type2, na.rm=TRUE)),]$row
          length(best_params)
          best Type2 <- dat result[which(dat result$Type2 ==</pre>
                                               max(dat_result$Type2, na.rm=TRUE)),]$Type2
In [37]: dfc01[best_params,]
          best_Type2
          A data.frame: 1 × 3
                  AST
                        CK Daysrec
                 <dbl> <dbl>
                               <dbl>
           3176
                 0.33
                       0.51
                                0.16
```

```
paste("Start time: ", start, sep="")
          dat_result <- gridSearch06(seed_vector, dat, dfc01)</pre>
          stop <- Sys.time()</pre>
          round(stop - start, 2)
          # Time difference of 41 mins
          'Start time: 2021-07-22 15:14:02'
          Time difference of 40.81 mins
In [39]: best params <- dat result[which(dat result$Type2 ==</pre>
                                               max(dat_result$Type2, na.rm=TRUE)),]$row
          length(best_params)
          best_Type2 <- dat_result[which(dat_result$Type2 ==</pre>
                                               max(dat_result$Type2, na.rm=TRUE)),]$Type2
          1
In [40]: dfc01[best_params,]
                                   CK
                         AST
                                          Daysrec
          # 25016
                        0.20
                                 0.51
                                              0.29
          best_Type2
          # 0.6634
          A data.frame: 1 × 3
                   AST
                         CK Daysrec
                 <dbl> <dbl>
                               <dbl>
           25016
                   0.2
                        0.51
                                 0.29
          0.66336
In [41]: # Compare the above weights with no weights.
          lst <- vector("list", length= 3)</pre>
          names(lst) <- c("AST","CK","Daysrec")</pre>
          lst[[1]] \leftarrow c(0.20)
          lst[[2]] \leftarrow c(0.51)
          lst[[3]] \leftarrow c(0.29)
          start <- Sys.time()</pre>
          dfc02 <- generate_combs(lst)</pre>
          stop <- Sys.time()</pre>
          # round(stop - start, 2)
           (dfc02 \leftarrow rbind(dfc02, rep(1, 3)))
          A data.frame: 2 x 3
            AST
                   CK Daysrec
```

```
start <- Sys.time()</pre>
         paste("Start time: ", start, sep="")
         dat_result <- gridSearch06(seed_vector, dat, dfc02)</pre>
         stop <- Sys.time()</pre>
         round(stop - start, 2)
         # Time difference of 3 mins
          'Start time: 2021-07-22 15:59:27'
         Time difference of 3.09 mins
In [43]: # The weights certainly make a difference, but not as
         # much of a difference for the Type2 score as I had
         # expected. The delta is only 0.008 (p-value of
         # around 1.7e-14 if we assume a standard deviation of
         # 0.0538 for each distribution; the sd value is estimated
         # from the comparative cross-val scores that follow).
         dat_result
         A data.frame: 2 ×
```

2
row Type2
<chr> <dbl> 1 0.66142
2 0.65326

Compare current weights with those from a previous version of this notebo

In a previous version of this notebook the seeds used were the same, but the rows of dat started in a difference up with a very different set of weights. Different combinations of weights can result in very similar Type 2 scores. Here I compare the previous weights with the current weights, but using 500 seeds. (Also differences in these 2 sets of weights says something about the adequacy of averaging over only 55 folds:

```
In [44]: lst <- vector("list", length= 3)
    names(lst) <- c("AST", "CK", "Daysrec")

lst[[1]] <- c(0.20)
lst[[2]] <- c(0.51)
lst[[3]] <- c(0.29)

start <- Sys.time()
dfc03 <- generate_combs(lst)
stop <- Sys.time()
# round(stop - start, 2)

dfc03 <- rbind(dfc03, c(0.59, 0.27, 0.14))</pre>
In []: set.seed(1233)
seed_vector <- sample(1:9999, 500, replace=FALSE)
```

Comparative cross-val scores for base k-means model with w

```
In [48]: # This function is called by get_cvInfo. It returns a vector
          # of scores: f-score, accuracy, Type2, false negatives, and
          # false positives, in the stated order. The scores are for
          # the base k-means model with weights.
          get_cvScores_kmBase <- function(traindat, valdat) {</pre>
              #################################
              # Transform and scale training set data for the
              # k-means model.
              traindat$AST <- log(traindat$AST)</pre>
              traindat$CK <- log(traindat$CK)</pre>
              traindat$Daysrec <- sqrt(traindat$Daysrec)</pre>
              traindat_scaled <- scale(traindat[, -1], center=TRUE, scale=TRUE)</pre>
              centers <- attr(traindat_scaled, "scaled:center")</pre>
              scales <- attr(traindat scaled, "scaled:scale")</pre>
              ###############################
              # Apply weights to traindat.
              wghts <- c(0.20, 0.51, 0.29)^{\circ}0.5
              names(wghts) <- cols <- c("AST", "CK", "Daysrec")</pre>
              df2 <- t(t(traindat_scaled[, cols]) * as.numeric(wghts[cols]))</pre>
              traindat_wghts <- as.data.frame(df2, row.names= rownames(traindat))</pre>
              colnames(traindat_wghts) <- cols</pre>
              ###############################
              # Prepare valdat.
              # Transform and scale valdat.
              valdat$AST <- log(valdat$AST)</pre>
              valdat$CK <- log(valdat$CK)</pre>
              valdat$Daysrec <- sqrt(valdat$Daysrec)</pre>
              valdat_scaled <- scale(valdat[, -1], center=centers, scale=scales)</pre>
              valdat_scaled <- as.data.frame(valdat_scaled, row.names=rownames(valdat))</pre>
              # Apply weights to valdat.
              df2 <- t(t(valdat_scaled[, cols]) * as.numeric(wghts[cols]))</pre>
```

```
colnames(dfout) <- c("Outcome", "cluster")</pre>
              c1_to_Outcome1 <- c1_toLevel_1(dfout)</pre>
              ###############################
              # Apply the k-means model to valdat wghts.
              # Each element of the following list is a row of valdat wghts.
              valdat_asList <- split(valdat_wghts[, colnames(kmod$centers)],</pre>
                                        seq(nrow(valdat_wghts)))
              ctr_list <- vector("list", length= nrow(valdat))</pre>
              for(i in 1:nrow(valdat)) {
                   ctr_list[[i]] <- kmod$centers</pre>
              names(ctr_list) <- rownames(valdat)</pre>
              # Get the predictions for the validation set.
              cluster_assgns <- mcmapply(getCluster, valdat_asList, ctr_list,</pre>
                                            SIMPLIFY=TRUE, mc.cores=6)
              valdat_wghts$cluster <- as.numeric(cluster_assgns)</pre>
              valdat_wghts$pred_Outcome <- NA</pre>
              if(c1_to_Outcome1) {
                   valdat_wghts[which(valdat_wghts$cluster==1),]$pred_Outcome <- 1</pre>
                   valdat_wghts[which(valdat_wghts$cluster==2),]$pred_Outcome <- 0</pre>
              } else {
                   valdat_wghts[which(valdat_wghts$cluster==1),]$pred_Outcome <- 0</pre>
                   valdat_wghts[which(valdat_wghts$cluster==2),]$pred_Outcome <- 1</pre>
              }
              # Generate confusion matrix for the k-means clusters and
              # the corresponding f-score.
              preds <- as.factor(valdat wghts$pred Outcome)</pre>
              names(preds) <- rownames(valdat)</pre>
              ans <- get_confusion(preds, valdat[, "Outcome", drop=FALSE])</pre>
              mat <- as.matrix(ans[[1]])</pre>
              mat[is.na(mat)] <- 0</pre>
              fscore <- round(as.numeric(ans[[2]]), 4)</pre>
              acc <- round(sum(diag(mat))/floor(sum(mat)), 4)</pre>
              type2 <- round((0.4 * acc + 0.6 * ans[[2]]), 4)
              FN <- as.numeric(mat[2,1])</pre>
              FP <- as.numeric(mat[1,2])</pre>
              return(c(fscore,acc,type2,FN,FP))
          }
In [50]: # Get scores for the base k-means model (with weights).
          set.seed(1913)
          seed_vector <- sample(1:9999, 1000, replace=FALSE)</pre>
          start <- Sys.time()</pre>
          dat_result <- get_cvInfo(seed_vector, dat)</pre>
          stop <- Sys.time()</pre>
          round(stop - start. 2)
```

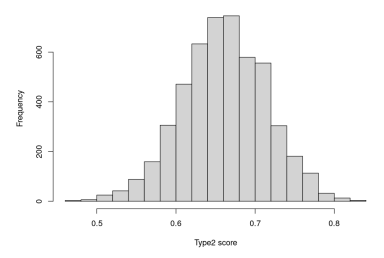
row.names=rownames(traindat))

```
A data.frame: 6 x 5
```

```
FP
                  fscore
                          Acc Type2
                                        FΝ
                   <dbl> <dbl>
                               <dbl> <dbl>
                                           <dbl>
           4782--1 0.5806 0.6750 0.6184
           4782--2 0.7018 0.7875 0.7361
                                         4
                                              13
           4782--3 0.6582 0.6625 0.6599
                                              17
           4782--4 0.6667 0.7000 0.6800
                                              17
           4782--5 0.5806 0.6750 0.6184
                                              18
           In [52]: | fscore mean <- round(mean(dat result$fscore), 4)</pre>
          fscore_sd <- round(sd(dat_result$fscore), 4)</pre>
          paste0("fscore mean: ", as.character(fscore_mean))
          # 0.6348
          paste0("fscore StdDev: ", as.character(fscore_sd))
          # 0.0620
          summary(dat_result$fscore)
          'fscore mean: 0.6348'
          'fscore StdDev: 0.062'
             Min. 1st Qu. Median
                                        Mean 3rd Qu.
                                                          Max.
                                       0.635 0.677
            0.385 0.594
                              0.636
                                                         0.824
In [53]: Acc_mean <- round(mean(dat_result$Acc), 4)</pre>
          Acc_sd <- round(sd(dat_result$Acc), 4)
paste0("accuracy mean: ", as.character(Acc_mean))</pre>
          paste0("accuracy StdDev: ", as.character(Acc_sd))
          # 0.0462
          summary(dat_result$Acc)
          'accuracy mean: 0.7004'
          'accuracy StdDev: 0.0462'
             Min. 1st Qu.
                             Median
                                        Mean 3rd Qu.
                                                          Max.
                                       0.700 0.738
            0.525
                    0.675
                              0.700
                                                         0.863
In [54]: Type2_mean <- round(mean(dat_result$Type2), 4)</pre>
          Type2_sd <- round(sd(dat_result$Type2), 4)</pre>
          paste0("Type2 mean: ", as.character(Type2_mean))
          # 0.6610
          paste0("Type2 StdDev: ", as.character(Type2_sd))
          # 0.0532
          summary(dat_result$Type2)
```

```
options(repr.plot.width= 8, repr.plot.height= 6)
hist(dat_result$Type2, breaks=14, xlab="Type2 score",
    main="Distribution of Type2 scores for base k-means model")
```

Distribution of Type2 scores for base k-means model



```
In [57]: FN_mean <- round(mean(dat_result$FN), 4)
FN_sd <- round(sd(dat_result$FN), 4)
paste0("FN mean: ", as.character(FN_mean))
# 8.72
paste0("FN StdDev: ", as.character(FN_sd))
# 2.46
""
summary(dat_result$FN)

'FN mean: 8.7152'
'FN StdDev: 2.4557'
"</pre>
```

Min. 1st Qu. Median Mean 3rd Qu. Max. 2.00 7.00 9.00 8.72 10.00 18.00

```
In [58]: FP_mean <- round(mean(dat_result$FP), 4)
    FP_sd <- round(sd(dat_result$FP), 4)
    paste0("FP mean: ", as.character(FP_mean))
# 15.25
    paste0("FP StdDev: ", as.character(FP_sd))
# 3.14
""
    summary(dat_result$FP)</pre>
```

'FP mean: 15.2548'
'FP StdDev: 3.1381'

The k-means model with weights has an f-score of 0.6348. svm02 has the next largest f-score: 0.6320. (hypothesis that 0.6348 > 0.6320 is 0.0124.) In terms of the f-score, then, the k-means model with weights the other three models. Its average number of false negatives is the lowest; but its average number of fal highest.

The k-means model with weights has an accuracy score of 0.7004. This score is lower than those for g00 svm02. svm02 has the next lowest accuracy score of 0.7030.

Overall, the k-means model with weights has a performance on par with those of our other three models: and svm02.

In Section 4 I construct a hybrid k-means model which out-performs all of the models that we have review

Section 4: k-means hybrid model

Add a predicted Outcome column (called "prob01") to the dataframe that k-means is applied to. See if thi column improves the clustering solution. This approach further leverages supervised learning to improve the unsupervised k-means algorithm. It would seem that in order for the prob01 column to boost the perfik-means model, it will need to not only (a) bring in new information; but also (b) add to, rather than subtrainformation already found in the other variables. In other words, the new information will need to complen conflict with, that found in the other variables.

Even though g03, svm02, and gbclf_best are nearly equal as classifiers, the differences between them a matters greatly which model we rely on to create a hybrid k-means model. In previous work I had first che because it has the highest Type2 score; but this choice weakened the performance of the hybrid model. I hybrid model with both svm02 and gbclf_best informing the k-means algorithm. This led to an even worse Next I chose the model (in this case, g03) with the best accuracy score rather than the best Type2 score. produced a better model. The approach is presented in what follows.

* * * * *

```
# been taken in the calling function.
cols <- names(wghts)</pre>
df2 <- t(t(traindat_scaled[, cols]) * as.numeric(wghts[cols]))</pre>
traindat_wghts <- as.data.frame(df2, row.names=rownames(traindat))</pre>
colnames(traindat_wghts) <- cols</pre>
###############################
# Prepare valdat.
# Compute the prob01 column.
preds01_b <- predict(g03mod, newdata=valdat)</pre>
valdat$prob01 <- as.numeric(preds01 b)</pre>
# Transform and scale valdat.
valdat$AST <- log(valdat$AST)</pre>
valdat$CK <- log(valdat$CK)</pre>
valdat$Daysrec <- sqrt(valdat$Daysrec)</pre>
valdat scaled <- scale(valdat[, -1], center=centers, scale=scales)</pre>
valdat scaled <- as.data.frame(valdat scaled, row.names=rownames(valdat))</pre>
# Apply weights to valdat.
df2 <- t(t(valdat_scaled[, cols]) * as.numeric(wghts[cols]))</pre>
valdat_wghts <- as.data.frame(df2, row.names=rownames(valdat))</pre>
colnames(valdat_wghts) <- cols</pre>
###############################
# Construct k-means model.
kmod <- suppressWarnings(kmeans(traindat_wghts, 2, iter.max = 50, nstart=</pre>
# See how the clusters are associated with Outcome.
dfout <- as.data.frame(cbind(traindat$Outcome, kmod$cluster),</pre>
                         row.names=rownames(traindat))
colnames(dfout) <- c("Outcome", "cluster")</pre>
c1_to_Outcome1 <- c1_toLevel_1(dfout)</pre>
##################################
# Apply the k-means model to valdat wghts.
# Each element of the following list is a row of valdat_wghts.
valdat_asList <- split(valdat_wghts[, colnames(kmod$centers)],</pre>
                         seq(nrow(valdat_wghts)))
ctr list <- vector("list", length= nrow(valdat))</pre>
for(i in 1:nrow(valdat)) {
    ctr_list[[i]] <- kmod$centers</pre>
names(ctr_list) <- rownames(valdat)</pre>
# Get the predictions for the validation set.
cluster_assgns <- mcmapply(getCluster, valdat_asList, ctr_list,</pre>
                             SIMPLIFY=TRUE, mc.cores=6)
valdat vahtatalvatas -
```

```
# Generate confusion matrix for the k-means clusters and
             # the corresponding f-score.
             preds <- as.factor(valdat_wghts$pred_Outcome)</pre>
             names(preds) <- rownames(valdat)</pre>
             ans <- get_confusion(preds, valdat[, "Outcome", drop=FALSE])</pre>
             # The result returned is a Type2 score (which is a mixture
             # of accuracy and f-score).
             mat <- as.matrix(ans[[1]])</pre>
             mat[is.na(mat)] <- 0</pre>
             percent_correct <- sum(diag(mat))/floor(sum(mat))</pre>
              result <- round((0.4 * percent\_correct + 0.6 * ans[[2]]), 6)
              return(result)
In [15]: # There are 4 parameter lists to work with. The best
         # approach, perhaps, is to start by exploring the
         # region around the space where all parameters have an
         # equal weight---in this case, a weight of 0.25.
         lst <- vector("list", length= 4)</pre>
         names(lst) <- c("AST","CK","Daysrec","prob01")</pre>
         lst[[1]] <- lst[[2]] <- lst[[3]] <- lst[[4]] <- seq(0.13, 0.37, by=0.02)
         start <- Sys.time()</pre>
         dfc01 <- generate_combs(lst)</pre>
         stop <- Sys.time()</pre>
         # round(stop - start, 2)
         dim(dfc01)
         # 1469
          1469 · 4
 In []: # Test on a sample of 10.
         set.seed(42)
         smp <- sample(rownames(dfc01), 10, replace=FALSE)</pre>
         tst_params <- dfc01[smp,]</pre>
         head(tst params)
 In [ ]: # Find the best weights of those in tst params.
         set.seed(1233)
         seed vector <- sample(1:9999, 10, replace=FALSE)</pre>
         start <- Sys.time()</pre>
         dat_result <- gridSearch06(seed_vector, dat, tst_params)</pre>
         stop <- Sys.time()</pre>
         round(stop - start, 2)
         # Time difference of 19 secs (for 10 rows, 10 seeds)
 In [ ]: best_params <- dat_result[which(dat_result$Type2 ==</pre>
                                           max(dat_result$Type2, na.rm=TRUE)),]$row
         length(best_params)
```

```
set.seed(1233)
          seed_vector <- sample(1:9999, 15, replace=FALSE)</pre>
          start <- Sys.time()</pre>
          paste("Start time: ", start, sep="")
          dat_result <- gridSearch06(seed_vector, dat, dfc01)</pre>
          stop <- Sys.time()</pre>
          round(stop - start, 2)
          # Time difference of 1.27 hours
          'Start time: 2021-07-23 10:18:29'
          Time difference of 1.27 hours
In [21]: best params <- dat result[which(dat result$Type2 ==</pre>
                                               max(dat result$Type2, na.rm=TRUE)),]$row
          length(best params)
          best_Type2 <- dat_result[which(dat_result$Type2 ==</pre>
                                               max(dat_result$Type2, na.rm=TRUE)),]$Type2
           1
In [22]: dfc01[best_params,]
                                   CK
                         AST
                                          Daysrec
                                                        prob01
          # 16213
                                 0.37
                                              0.21
                        0.15
                                                          0.27
          best_Type2
          # 0.6649
          A data.frame: 1 × 4
                  AST
                         CK Daysrec prob01
                 <dbl> <dbl>
                               <dbl>
                                      <dbl>
           16213 0.15
                        0.37
                                0.21
                                       0.27
          0.66489
In [27]: # Refine the search.
          lst <- vector("list", length= 4)</pre>
          names(lst) <- c("AST", "CK", "Daysrec", "prob01")</pre>
          lst[[1]] \leftarrow seq(0.09, 0.15, by= 0.02)
          lst[[2]] \leftarrow seq(0.37, 0.45, by= 0.01)
          lst[[3]] \leftarrow seq(0.11, 0.23, by= 0.02)
          lst[[4]] \leftarrow seq(0.25, 0.35, by=0.01)
          start <- Sys.time()</pre>
          dfc02 <- generate_combs(lst)</pre>
          stop <- Sys.time()</pre>
          # round(stop - start, 2)
          dim(dfc02)
```

```
In [29]: dat_result_dfc01 <- dat_result</pre>
In [30]: # Find the best weights of those in dfc02 (173 rows,
          # 15 seeds, 5 folds).
          set.seed(1233)
          seed_vector <- sample(1:9999, 15, replace=FALSE)</pre>
          start <- Sys.time()</pre>
          paste("Start time: ", start, sep="")
          dat_result <- gridSearch06(seed_vector, dat, dfc02)</pre>
          stop <- Sys.time()</pre>
           round(stop - start, 2)
          # Time difference of 9.48 mins
          'Start time: 2021-07-23 12:45:58'
          Time difference of 9.48 mins
In [31]: best_params <- dat_result[which(dat_result$Type2 ==</pre>
                                               max(dat result$Type2, na.rm=TRUE)),]$row
          length(best_params)
          best_Type2 <- dat_result[which(dat_result$Type2 ==</pre>
                                               max(dat result$Type2, na.rm=TRUE)),]$Type2
In [32]: dfc02[best params,]
                         AST
                                    CK
                                           Daysrec
                                                         prob01
          # 660
                        0.15
                                 0.39
                                              0.19
                                                           0.27
          best_Type2
          # 0.6655
          A data.frame: 1 × 4
                 AST
                        CK Daysrec prob01
                <dbl> <dbl>
                              <dbl>
                                     <dbl>
           660 0.15
                                      0.27
                      0.39
                               0.19
          0.66546
In [33]: # Refine the search.
          lst <- vector("list", length= 4)
names(lst) <- c("AST","CK","Daysrec","prob01")</pre>
          lst[[1]] \leftarrow seq(0.13, 0.17, by= 0.01)
          lst[[2]] \leftarrow seq(0.37, 0.43, by= 0.01)
          lst[[3]] \leftarrow seq(0.15, 0.21, by= 0.01)
          lst[[4]] \leftarrow seq(0.25, 0.31, by=0.01)
          start <- Svs time()</pre>
```

```
In [34]: dat_result_dfc02 <- dat_result</pre>
In [35]: # Find the best weights of those in dfc02 (170 rows,
          # 15 seeds, 5 folds).
          set.seed(1233)
          seed_vector <- sample(1:9999, 15, replace=FALSE)</pre>
          start <- Sys.time()</pre>
          paste("Start time: ", start, sep="")
          dat_result <- gridSearch06(seed_vector, dat, dfc03)</pre>
          stop <- Sys.time()</pre>
          round(stop - start, 2)
          # Time difference of 9 mins
          'Start time: 2021-07-23 13:01:03'
          Time difference of 9.06 mins
In [36]: best_params <- dat_result[which(dat_result$Type2 ==</pre>
                                             max(dat result$Type2, na.rm=TRUE)),]$row
          length(best_params)
          best_Type2 <- dat_result[which(dat_result$Type2 ==</pre>
                                             max(dat result$Type2, na.rm=TRUE)),]$Type2
In [37]: dfc03[best params,]
                                  CK
                                                      prob01
                        AST
                                        Daysrec
          # 647
                       0.14
                             0.40
                                            0.19
                                                        0.27
          best_Type2
          # 0.6655
          A data.frame: 1 × 4
                AST
                      CK Daysrec prob01
               <dbl> <dbl>
                            <dbl>
                                   <dbl>
           647 0.14
                             0.19
                                    0.27
                       0.4
          0.66551
```

Summary info for hybrid model with prob01 constructed from

First change get_cvInfo to call the function in the next cell.

```
In [45]: # This function is called by get_cvInfo. It returns a vector
# of scores: f-score, accuracy, Type2, false negatives, and
# false positives, in the stated order. The scores are for
# the hybrid k-means model with the prob01 column.
```

```
traindat$AST <- log(traindat$AST)</pre>
traindat$CK <- log(traindat$CK)</pre>
traindat$Daysrec <- sqrt(traindat$Daysrec)</pre>
traindat_scaled <- scale(traindat[, -1], center=TRUE, scale=TRUE)</pre>
centers <- attr(traindat_scaled, "scaled:center")
scales <- attr(traindat_scaled, "scaled:scale")</pre>
##################################
# Apply weights to traindat.
wghts <- c(0.14, 0.40, 0.19, 0.27)^{\circ}0.5
names(wghts) <- cols <- c("AST","CK","Daysrec","prob01")
df2 <- t(t(traindat_scaled[, cols]) * as.numeric(wghts[cols]))</pre>
traindat_wghts <- as.data.frame(df2, row.names=rownames(traindat))</pre>
colnames(traindat_wghts) <- cols</pre>
###############################
# Prepare valdat.
# Compute the prob01 column.
preds01_b <- predict(g03mod, newdata=valdat)</pre>
valdat$prob01 <- as.numeric(preds01_b)</pre>
# Transform and scale valdat.
valdat$AST <- log(valdat$AST)</pre>
valdat$CK <- log(valdat$CK)</pre>
valdat$Daysrec <- sqrt(valdat$Daysrec)</pre>
valdat_scaled <- scale(valdat[, -1], center=centers, scale=scales)</pre>
valdat_scaled <- as.data.frame(valdat_scaled, row.names=rownames(valdat))</pre>
# Apply weights to valdat.
df2 <- t(t(valdat_scaled[, cols]) * as.numeric(wghts[cols]))</pre>
valdat wghts <- as.data.frame(df2, row.names=rownames(valdat))</pre>
colnames(valdat wghts) <- cols</pre>
###############################
# Construct k-means model.
kmod <- suppressWarnings(kmeans(traindat wghts, 2, iter.max = 50, nstart=
# See how the clusters are associated with Outcome.
dfout <- as.data.frame(cbind(traindat$Outcome, kmod$cluster),</pre>
                           row.names=rownames(traindat))
colnames(dfout) <- c("Outcome", "cluster")</pre>
c1_to_Outcome1 <- c1_toLevel_1(dfout)</pre>
###############################
# Apply the k-means model to valdat_wghts.
# Each element of the following list is a row of valdat_wghts.
valdat_asList <- split(valdat_wghts[, colnames(kmod$centers)],</pre>
                           seq(nrow(valdat wghts)))
ctr_list <- vector("list", length= nrow(valdat))
far(i in larger(valdat));</pre>
```

```
valdat_wghts$cluster <- as.numeric(cluster_assgns)</pre>
              valdat_wghts$pred_Outcome <- NA</pre>
              if(c1_to_Outcome1) {
                   valdat_wghts[which(valdat_wghts$cluster==1),]$pred_Outcome <- 1</pre>
                   valdat_wghts[which(valdat_wghts$cluster==2),]$pred_Outcome <- 0</pre>
                   valdat_wghts[which(valdat_wghts$cluster==1),]$pred_Outcome <- 0</pre>
                   valdat_wghts[which(valdat_wghts$cluster==2),]$pred_Outcome <- 1</pre>
              }
              # Generate confusion matrix for the k-means clusters and
              # the corresponding f-score.
              preds <- as.factor(valdat wghts$pred Outcome)</pre>
              names(preds) <- rownames(valdat)</pre>
              ans <- get_confusion(preds, valdat[, "Outcome", drop=FALSE])</pre>
              mat <- as.matrix(ans[[1]])</pre>
              mat[is.na(mat)] <- 0</pre>
              fscore <- round(as.numeric(ans[[2]]), 4)</pre>
              acc <- round(sum(diag(mat))/floor(sum(mat)), 4)</pre>
              type2 <- round((0.4 * acc + 0.6 * ans[[2]]), 4)
              FN <- as.numeric(mat[2,1])</pre>
              FP <- as.numeric(mat[1,2])</pre>
              return(c(fscore,acc,type2,FN,FP))
In [46]: # Get scores for the hybrid model with prob01 constructed
          # from the g03 logistic model.
          set.seed(1913)
          seed_vector <- sample(1:9999, 1000, replace=FALSE)</pre>
          start <- Sys.time()</pre>
          dat_result <- get_cvInfo(seed_vector, dat)</pre>
          stop <- Sys.time()</pre>
          round(stop - start, 2)
          # Time difference of 3.53 mins
```

Time difference of $3.4\ \mathrm{mins}$

```
In [47]: dim(dat_result)
head(dat_result)
```

5000 · 5

A data.frame: 6 × 5

	fscore	Acc	Type2	FN	FP	
	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	
47821	0.4800	0.6750	0.5580	20	6	
47822	0.6531	0.7875	0.7069	8	9	
47823	0.6571	0.7000	0.6743	13	11	
47824	0.6667	0.7375	0.6950	10	11	
47825	0.6182	0.7375	0.6659	9	12	

0.456

0.625

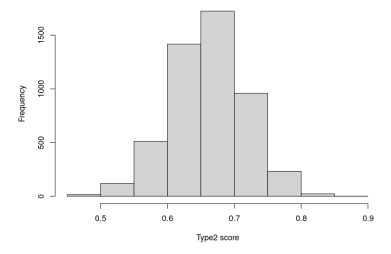
0.662

0.662 0.700

0.860

```
'fscore mean: 0.6163'
          'fscore StdDev: 0.0676'
             Min. 1st Qu. Median
                                      Mean 3rd Qu.
                                                       Max.
            0.323 0.571
                            0.618
                                      0.616 0.667
                                                       0.842
In [49]: Acc_mean <- round(mean(dat_result$Acc), 4)</pre>
          Acc_sd <- round(sd(dat_result$Acc), 4)</pre>
          paste0("accuracy mean: ", as.character(Acc_mean))
          paste0("accuracy StdDev: ", as.character(Acc_sd))
          # 0.0443
          summary(dat_result$Acc)
          'accuracy mean: 0.7298'
          'accuracy StdDev: 0.0443'
             Min. 1st Qu. Median
                                      Mean 3rd Qu.
                                                       Max.
            0.575 0.700
                             0.725
                                      0.730 0.762
                                                       0.887
In [50]: Type2_mean <- round(mean(dat_result$Type2), 4)</pre>
          Type2_sd <- round(sd(dat_result$Type2), 4)</pre>
          paste0("Type2 mean: ", as.character(Type2_mean))
          paste0("Type2 StdDev: ", as.character(Type2_sd))
          # 0.0556
          summary(dat_result$Type2)
          'Type2 mean: 0.6617'
          'Type2 StdDev: 0.0556'
                                      Mean 3rd Qu.
             Min. 1st Qu. Median
                                                       Max.
```

Distribution of Type2 scores for hybrid model w/ prob01 from g03



```
In [52]: FN_mean <- round(mean(dat_result$FN), 4)
    FN_sd <- round(sd(dat_result$FN), 4)
    paste0("FN mean: ", as.character(FN_mean))
# 12.22
    paste0("FN StdDev: ", as.character(FN_sd))
# 3.06
""
summary(dat_result$FN)</pre>
```

'FN mean: 12.2196'

'FN StdDev: 3.0643'

"

Min. 1st Qu. Median Mean 3rd Qu. Max. 3.0 10.0 12.0 12.2 14.0 23.0

```
In [53]: FP_mean <- round(mean(dat_result$FP), 4)
    FP_sd <- round(sd(dat_result$FP), 4)
    paste0("FP mean: ", as.character(FP_mean))
# 9.40
    paste0("FP StdDev: ", as.character(FP_sd))
# 3.00
""
summary(dat_result$FP)</pre>
```

'FP mean: 9.3952'

'FP StdDev: 3.0007'

```
header=TRUE, row.names=1)
           dim(results)
            5 · 10
 In [ ]: #&* Bookmark
In [55]: # The following table is a summary of all the
           # results from Sections 3 and 4. Recall that
           # the Type2 score is 60% f-score and 40% accuracy.
           results
           A data frame: 5 x 10
                                                                                          FP FP_sd
                      fscore
                             fscore_sd accuracy acc_sd Type2 Type2_sd
                                                                            FN FN_sd
                       <dbl>
                                 <dbl>
                                          <dbl>
                                                  <dbl>
                                                         <dbl>
                                                                   <dbl>
                                                                         <dbl>
                                                                                 <dbl>
                                                                                       <dbl>
                                                                                              <dbl>
                 g03 0.6183
                                0.0646
                                          0.7167
                                                 0.0448 0.6577
                                                                  0.0543
                                                                          11.24
                                                                                  3.04
                                                                                       11.43
                                                                                               3.34
                gbclf 0.6051
                                0.0665
                                          0.7135
                                                 0.0448
                                                        0.6485
                                                                  0.0552
                                                                          12.03
                                                                                  3.23
                                                                                       10.89
                                                                                               3.29
               svm02 0.6320
                                0.0628
                                          0.7030
                                                 0.0472 0.6604
                                                                  0.0542
                                                                          14.53
                                                                                  3.63
                                                                                        9.23
                                                                                               3.16
            km_wghts 0.6348
                                          0.7004
                                                 0.0462
                                                                  0.0532
                                                                           8.72
                                0.0620
                                                        0.6610
                                                                                  2.46
                                                                                       15.25
                                                                                               3.14
            kmp1_g03 0.6163
                                0.0676
                                          0.7298
                                                 0.0443 0.6617
                                                                  0.0556
                                                                          12.22
                                                                                  3.06
                                                                                        9.40
                                                                                               3.00
 In [ ]:
```

Final Comments for Section 4

Of the entries in the above table, the hybrid k-means model has the best accuracy score (0.7298) and the (0.6617). The next best non-k-means Type2 score is for svm02 (0.6604). The p-value for the hypothesis (0.12, so the difference we see is not statistically significant. However, the hybrid model is clearly a better because it has a much higher accuracy score (p-value for the delta is < 2e-16); on average, the hybrid m significantly fewer false negatives than svm02 and nearly the same average number of false positives.

The hybrid model also outperforms g03, the model with the next highest accuracy score (p-value for the On average, the hybrid model has significantly fewer false positives than g03.

The 2 k-means models in the table are about equal in overall performance. The hybrid model has signific positives, but significantly more false negatives. The hybrid model has a significantly lower f-score, but it with its significantly higher accuracy score. If the k-means algorithm was less suited to the cow data, km_ be nearly so equal in performance to kmp1_g03.

* * * * *

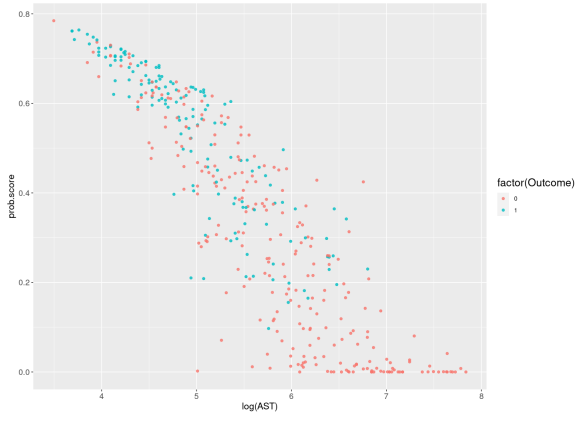
We can try to further boost the above hybrid model by adding a second probability column (prob02). In o in performance, the additional column will need to contain new information that complements what is alre other columns. If we generate prob02 using gbclf_best, the new hybrid model (at least in the testing I hav perform any better than kmp1_g03.

* * * * :

```
In [ ]:
```

Addendum: More plots showing separation of points Outcome level

Differences in AST and prob01, by Outcome level

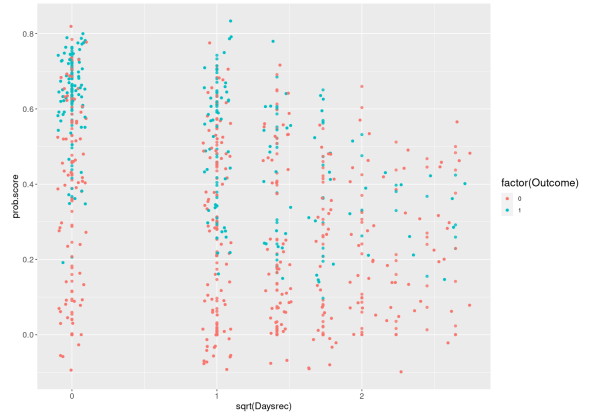


```
options(repr.plot.width= 12, repr.plot.height= 9)

p <- ggplot(dat_plot, aes(Daysrec, prob01, color= factor(Outcome))) +

geom_point(alpha= 0.8) + geom_jitter(width= 0.1, height= 0.1) +
 xlab("sqrt(Daysrec)") + ylab("prob.score") +
 ggtitle("Differences in Daysrec and prob01, by Outcome level") +
 theme(axis.text= element_text(size= 12)) +
 theme(axis.title= element_text(size= 14)) +
 theme(title= element_text(size= 16))
p</pre>
```

Differences in Daysrec and prob01, by Outcome level



```
In [15]: # Plot Daysrec vs. AST.

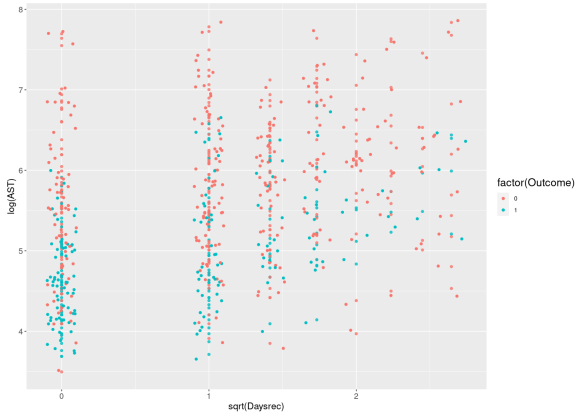
dat_plot <- newtrain[, c("Outcome", "Daysrec", "AST"), drop=FALSE]
dat_plot$Daysrec <- sqrt(dat_plot$Daysrec)
dat_plot$AST <- log(dat_plot$AST)

options(repr.plot.width= 12, repr.plot.height= 9)

p <- ggplot(dat_plot, aes(Daysrec, AST, color= factor(Outcome))) +

geom_point(alpha= 0.8) + geom_jitter(width= 0.1, height= 0.1) +
    xlab("sqrt(Daysrec)") + ylab("log(AST)") +
    ggtitle("Differences in Daysrec and AST, by Outcome level") +
    theme(axis.text= element_text(size= 12)) +
    theme(axis.title= element_text(size= 14)) +
    theme(title= element_text(size= 16))
</pre>
```

Differences in Daysrec and AST, by Outcome level



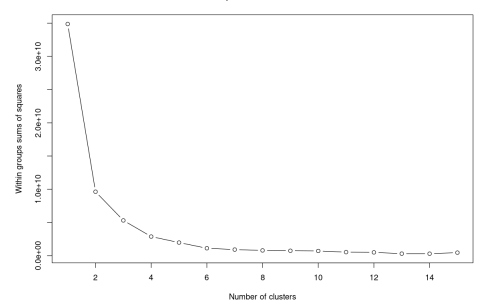
Postscript: Using tot.withinss to search for weights

When the k-means algorithm is applied to the data on downer cows, we ask the algorithm to give us 2 cluster interested in only 2 clusters---distinguishing between the cows that survived and those that did not. B from the wss (within-group sum of squares) plot below, that a 2-cluster solution is optimal for our dataset interest in distinguishing between survivors and non-survivors. We might say that the dataset naturally le

.

```
In [17]: options(repr.plot.width= 10, repr.plot.height= 7)
# Remove the Outcome column from data before plotting.
wssplot(dat[, -1], title= "wss plot for the cow data")
```

wss plot for the cow data

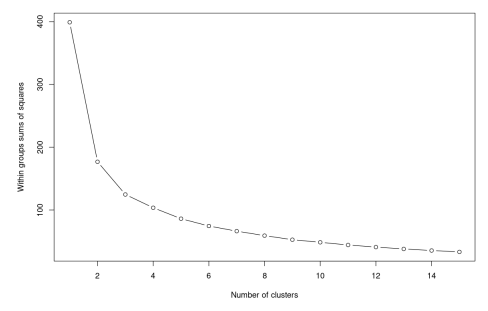


```
In []: ### COMMENT (on above graph):
    # The largest gain we see in the reduction of within-group
    # sum of squares occurs when we have 2 clusters. This
    # strongly suggests a 2-cluster solution to the 400
    # observations in dat. The additional gain we would
    # get from a 3-cluster solution is negligible relative to
    # the gain we see from a 2-cluster solution. Thus, a
```

```
In [18]: # Transform the data as if modeling for k-means.
           df <- dat
           g03 <- glm(Outcome ~ Daysrec + CK + I(log(AST)),
                                data= df, family= binomial, singular.ok=TRUE,
                                epsilon= 1e-7, maxit=50)
           df$prob01 <- as.numeric(g03$fitted)</pre>
           df$AST <- log(df$AST)</pre>
           df$CK <- log(df$CK)</pre>
           df$Daysrec <- sqrt(df$Daysrec)</pre>
           df_scaled <- scale(df[, -1], center=TRUE, scale=TRUE)</pre>
           df_scaled <- as.data.frame(cbind(df$Outcome, df_scaled),</pre>
                                          row.names=rownames(df))
           colnames(df_scaled) <- colnames(df)</pre>
           # The following weights are those found in the above
           # grid search for the hybrid model, kmp1_g03.
           wghts <- c(0.14, 0.40, 0.19, 0.27)^{\circ}0.5
           names(wghts) <- cols <- c("AST", "CK", "Daysrec", "prob01")</pre>
           df2 <- t(t(df_scaled[, cols]) * as.numeric(wghts[cols]))</pre>
           df_wghts <- cbind(as.numeric(df_scaled$Outcome), df2)</pre>
          df_wghts <- as.data.frame(df_wghts)
colnames(df_wghts) <- c("Outcome", cols)
rownames(df_wghts) <- rownames(df_scaled)</pre>
In [19]: options(repr.plot.width= 10, repr.plot.height= 7)
           # Remove the Outcome column from data before plotting.
```

wssplot(df_wghts[, -1], title= "wss plot: cow data, scaled, + prob01")

wss plot: cow data, scaled, + prob01



I want to see if this approach to finding weights works and, if so, whether it is a faster way to find optimal

* * * * *

It turns out that the weights we get using this approach depend very much on the scaling of the variables min-max scaling *after* centering and scaling each of the variables. The mins and maxs from the training standard for the scaling of the validation data. With this dataset, doing so will not get us usable weights.

```
In [21]: # Function for computing the tot.withinss for each set of
          # weights in df_params (a dataframe, each row of which is
          # a candidate set of weights). The optimal set of weights
          # will be the set that yields the smallest average (over
          # the folds) for tot.withinss.
          # This function is called from gridSearch07.
          get_tot.withinss_g03 <- function(traindat, valdat, wghts) {</pre>
              g03mod <- suppressWarnings(glm(Outcome ~ Daysrec + CK + I(log(AST)),</pre>
                              data= traindat, family= binomial, singular.ok=TRUE,
                              epsilon= 1e-7, maxit=50))
              traindat$prob01 <- as.numeric(g03mod$fitted)</pre>
              # Scale training set data. We need 'centers' and 'scales'
              # for scaling valdat.
              df <- scale(traindat[, -1], center=TRUE, scale=TRUE)</pre>
              centers <- attr(df, "scaled:center")
scales <- attr(df, "scaled:scale")</pre>
              df <- as.matrix(df)</pre>
              traindat_scaled <- apply(df, MARGIN=2, range01)</pre>
              colnames(traindat_scaled) <- colnames(traindat)[-1]</pre>
              rownames(traindat_scaled) <- rownames(traindat)</pre>
              # Get mins and maxs for scaling of valdat.
              # traindat_mins <- as.numeric(apply(df, MARGIN=2, min))</pre>
              # traindat_maxs <- as.numeric(apply(df, MARGIN=2, max))</pre>
              ###############################
              # Transform and scale valdat.
              valdat$prob01 <- as.numeric(predict(g03mod, newdata=valdat))</pre>
              df02 <- scale(valdat[, -1], center=centers, scale=scales)</pre>
              df02 <- as.matrix(df02)</pre>
              # df02_t <- t(as.matrix(df02))
              # df02_asList <- split(df02_t, seq(nrow(df02_t)))</pre>
              # names(df02_asList) <- colnames(valdat)[-1]</pre>
              # valdat_scaled <- mapply(range02, df02_asList, traindat_mins,</pre>
                                          traindat_maxs)
              valdat_scaled <- apply(df02, MARGIN=2, range01)</pre>
              # The next step is crucial.
              valdat scaled <- as.data.frame(valdat scaled, row.names=rownames(valdat))</pre>
              colnames(valdat_scaled) <- colnames(valdat)[-1]</pre>
              # Applv weights to valdat.
```

```
In [17]: # This grid search searches for the best set of weights to use
          # in our k-means clustering model. The best weights are those
          # which generalize best to the validation set. So we look for
          # the best cross-validation score.
          # Because our training set is so small---only 400 records---we
          # need to run the gridSearch over many seeds. Otherwise, we
          # will not get a meaningful result.
          gridSearch07 <- function(seed_vector, dat, df_params, folds=5) {</pre>
              datout <- rep(NA, 2*nrow(df params))</pre>
              dim(datout) <- c(nrow(df_params), 2)</pre>
              datout <- as.data.frame(datout)</pre>
              colnames(datout) <- c("row", "tot.withinss")</pre>
              datout$row <- params_rows <- rownames(df_params)</pre>
              # We want the sgrt of the weights.
              df params <- df params^0.5
              ################################
              # Partition the data into folds.
              segment_size <- round(dim(dat)[1]/folds)</pre>
              diff <- dim(dat)[1] - folds * segment_size</pre>
              last seg size <- segment size + diff</pre>
              segmentsv <- c(rep(segment size, (folds - 1)), last seg size)</pre>
              stopifnot(sum(segmentsv) == dim(dat)[1])
              # Create a dataframe, each row for a distinct seed.
              # Each column of the dataframe is for a distinct set
              # of weights. The entries in the cells are tot.withinss
              # scores.
              seedv_len <- length(seed_vector)</pre>
              df_scores <- rep(NA, seedv_len*nrow(df_params))</pre>
              dim(df_scores) <- c(seedv_len, nrow(df_params))</pre>
              df_scores <- as.data.frame(df_scores)</pre>
              colnames(df_scores) <- rownames(df_params)</pre>
              rownames(df_scores) <- as.character(seed_vector)</pre>
              for(h in 1:seedv_len) {
                  # shuffle dat
                  cur seed <- seed vector[h]</pre>
                  set.seed(cur_seed)
                  smp <- sample(rownames(dat), nrow(dat), replace= FALSE)</pre>
                  dat <- dat[smp,]</pre>
                  # Each element of row_list will be the rows we pick
                  # out for one of the folds. E.g., the first element
                  # of row_list will contain the rows we want for the
                  # first fold, the second element of row list will
                  # contain the rows we want for the second fold, and
                  # so forth.
                  row list <- vector("list", length=folds)</pre>
                  names(row_list) <- as.character(1:folds)</pre>
                  startpt <- 1
                  for(i in 1:folds) {
                       endpt <- startpt + segmentsv[i] - 1</pre>
                       # stopifnot(endpt <= nrow(dat))</pre>
```

```
for(j in 1:folds) {
                           testdat <- dat[row_list[[j]],]</pre>
                           traindat <- dat[which(!(rownames(dat) %in% rownames(testdat))</pre>
                           # stopifnot((length(rownames(traindat)) + length(rownames(tes
                           test_list[[j]] <- testdat</pre>
                           train_list[[j]] <- traindat</pre>
                       # When there are only 5 folds, only 5 cores get used.
                       scores <- mcmapply(get_tot.withinss_g03, train_list, test_list,</pre>
                                           MoreArgs= list(wghts=wghts),
                                           SIMPLIFY= TRUE, mc.cores=5)
                       # For the current seed, store the average of the tot.withinss
                       # scores, the average taken over the folds.
                       df_scores[as.character(cur_seed), cur_row] <- round(mean(scores),</pre>
                  } # end of for-loop, index i
              } ## end of for-loop, index h
              # Compute the average over the seeds of the tot.withinss scores
              # obtained for each set of parameters in df params.
              datout$tot.withinss <- round(apply(df_scores, MARGIN=2, mean), 5)</pre>
              return(datout)
In [18]: # There are 4 parameter lists to work with.
          lst <- vector("list", length= 4)</pre>
          names(lst) <- c("AST", "CK", "Daysrec", "prob01")</pre>
          lst[[1]] <- lst[[2]] <- lst[[3]] <- lst[[4]] <- seq(0.13, 0.37, by=0.02)
          start <- Sys.time()</pre>
          dfc01 <- generate_combs(lst)</pre>
          stop <- Sys.time()</pre>
          # round(stop - start, 2)
          dim(dfc01)
          # 1469
          1469 · 4
In [19]: # Test on a sample of 10.
          set.seed(42)
          smp <- sample(rownames(dfc01), 10, replace=FALSE)</pre>
          tst_params <- dfc01[smp,]</pre>
          # head(tst_params)
 In [ ]: # Find the best weights of those in tst_params.
          set.seed(1233)
          seed_vector <- sample(1:9999, 11, replace=FALSE)</pre>
          start <- Sys.time()</pre>
          dat_result <- gridSearch07(seed_vector, dat, tst_params)</pre>
```

```
In [ ]: dfc01[best_params,]
          best_tot.withinss
In [25]: # Find the best weights of those in dfc01 (1469 rows,
          # 15 seeds, 5 folds).
          set.seed(1233)
          seed_vector <- sample(1:9999, 15, replace=FALSE)</pre>
          start <- Sys.time()</pre>
          paste0("Start time: ", start)
          dat_result <- gridSearch07(seed_vector, dat, dfc01)</pre>
          stop <- Sys.time()</pre>
          round(stop - start, 2)
          # Time difference of 10.6 - 13.2 mins (= 0.341 secs/row)
          # This is 7X faster than the first approach which
          # used cross-validation with the Type2 score.
          'Start time: 2021-07-25 10:35:57'
          Time difference of 13.17 mins
In [26]: best params <- dat result[which(dat result$tot.withinss ==</pre>
                                             min(dat_result$tot.withinss, na.rm=TRUE)),]$r
          length(best params)
          best_tot.withinss <- round(dat_result[which(dat_result$tot.withinss ==</pre>
                                             min(dat_result$tot.withinss, na.rm=TRUE)),]$t
          1
In [27]: dfc01[best_params,]
                                  CK
                        AST
                                        Daysrec
                                                     prob01
          # 26521
                       0.13
                                0.37
                                            0.13
                                                        0.37
          best_tot.withinss
          # 1.42
          A data.frame: 1 × 4
                  AST
                        CK Daysrec prob01
                 <dbl> <dbl>
                              <dbl>
                                    <dbl>
           26521
                 0.13
                       0.37
                               0.13
                                      0.37
          1.4202
In [28]: # Refine the search.
          lst <- vector("list", length= 4)</pre>
          names(lst) <- c("AST", "CK", "Daysrec", "prob01")</pre>
```

```
dim(dfc02)
          # 1091
                       4
           1091 · 4
In [29]: # Find the best weights of those in dfc02 (1091 rows,
          # 15 seeds, 5 folds).
          set.seed(1233)
          seed vector <- sample(1:9999, 15, replace=FALSE)</pre>
          start <- Sys.time()</pre>
          paste("Start time: ", start, sep="")
          dat_result <- gridSearch07(seed_vector, dat, dfc02)</pre>
          stop <- Sys.time()</pre>
          round(stop - start, 2)
          # Time difference of 10.2 mins
          'Start time: 2021-07-25 10:52:39'
          Time difference of 10.2 mins
In [30]: best params <- dat result[which(dat result$tot.withinss ==</pre>
                                              min(dat_result$tot.withinss, na.rm=TRUE)),]$r
          length(best_params)
          best_tot.withinss <- round(dat_result[which(dat_result$tot.withinss ==</pre>
                                              min(dat_result$tot.withinss, na.rm=TRUE)),]$t
In [31]: dfc02[best params,]
                                  CK
                                                       prob01
                        AST
                                         Daysrec
          # 12717
                       0.04
                                0.48
                                             0.04
                                                         0.44
          best_tot.withinss
          # 0.92
          A data.frame: 1 × 4
                  AST
                         CK Daysrec prob01
                 <dbl> <dbl>
                              <dbl>
                                     <dbl>
           12717
                  0.04
                        0.48
                                0.04
                                      0.44
          0.9184
In [32]: # Of the 2 sets of weights identified, see which set does
          # better when we use cross-validation with the Type2 score.
          lst <- vector("list", length= 4)
names(lst) <- c("AST","CK","Daysrec","prob01")</pre>
          lst[[1]] <- 0.04
          lst[[2]] <- 0.48
```

A data.frame: 3 × 4

prob01	Daysrec	CK	AST
<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
0.44	0.04	0.48	0.04
0.37	0.13	0.37	0.13
0.27	0.19	0.40	0.14

```
In [33]: # Find the best weights of those in dfc02pt2
# using 500 seeds.

set.seed(1233)
seed_vector <- sample(1:9999, 500, replace=FALSE)

start <- Sys.time()
dat_result <- gridSearch06(seed_vector, dat, dfc02pt2)
stop <- Sys.time()
round(stop - start, 2)
# Time difference of 5.6 mins</pre>
```

Time difference of 5.59 mins

```
In [34]: datout <- cbind(dfc02pt2, dat_result$Type2)
   colnames(datout) <- c(colnames(dfc02pt2), "Type2")
   datout</pre>
```

A data.frame: 3 × 5

AST	СК	Daysrec	prob01	Type2
<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
0.04	0.48	0.04	0.44	0.66055
0.13	0.37	0.13	0.37	0.66094
0.14	0.40	0.19	0.27	0.66175

```
In [ ]: ### COMMENTS:
```

```
# The Type2 score of the weights found in Section 4 is the
# highest score. These are the weights used for the
# kmp1_g03 model. The next best weights are those first
# found using tot.withinss (2nd row in above table).

# Perhaps we can use tot.withinss for the first round or two
# of the search for best weights and then finish the search
# relying on the Type2 score. This is what I proceed to do
# in the cells that follow. With the restricted range I have
# chosen for dfc03, we should settle on nearly the same weights
# we found in Section 4 (3rd row in the above table).
```

```
In [35]: # Refine the search for weights, starting with the set
    # initially found using tot.withinss. Run cross-validation
    # using Type2 scores.
```

```
# 35
                     4
          35 · 4
In [36]: # Find the best weights of those in dfc02 (35 rows,
          # 51 seeds, 5 folds). Note that I am using 51 seeds
          # here rather than 11.
          set.seed(1233)
          seed vector <- sample(1:9999, 51, replace=FALSE)</pre>
          start <- Sys.time()</pre>
          paste("Start time: ", start, sep="")
          dat_result <- gridSearch06(seed_vector, dat, dfc03)</pre>
          stop <- Sys.time()</pre>
          round(stop - start, 2)
          # Time difference of 6.45 mins
          'Start time: 2021-07-25 11:20:25'
          Time difference of 6.45 mins
In [38]: best_params <- dat_result[which(dat_result$Type2 ==</pre>
                                             max(dat_result$Type2, na.rm=TRUE)),]$row
          length(best_params)
          best Type2 <- dat result[which(dat result$Type2 ==</pre>
                                             max(dat result$Type2, na.rm=TRUE)),]$Type2
In [39]: dfc03[best_params,]
                        AST
                                  CK
                                        Daysrec
                                                      prob01
          # 359
                       0.13
                                0.38
                                            0.12
                                                        0.37
          best_Type2
          # 0.6622
          A data.frame: 1 × 4
                AST
                      CK Daysrec prob01
               <dbl> <dbl>
                            <dbl>
                                   <dbl>
               0.13
                     0.38
                             0.12
                                    0.37
           359
          0.66219
```

Summary info-2 for hybrid model with prob01 constructed from

First change get_cvInfo to call the function in the next cell.

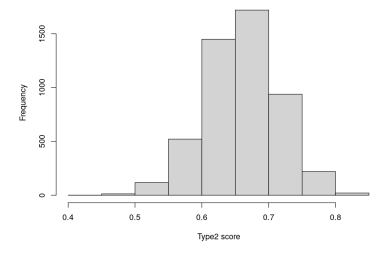
```
In [40]: # This function is called by get_cvInfo. It returns a vector # of scores: f-score. accuracy. Type2. false negatives. and
```

```
traindat$prob01 <- as.numeric(g03mod$fitted)</pre>
###############################
# Transform and scale training set data.
traindat$AST <- log(traindat$AST)</pre>
traindat$CK <- log(traindat$CK)</pre>
traindat$Daysrec <- sqrt(traindat$Daysrec)</pre>
traindat_scaled <- scale(traindat[, -1], center=TRUE, scale=TRUE)</pre>
centers <- attr(traindat_scaled, "scaled:center")</pre>
scales <- attr(traindat_scaled, "scaled:scale")</pre>
###################################
# Apply weights to traindat.
# These are the most recents weights we have found.
wghts \leftarrow c(0.13, 0.38, 0.12, 0.37)^0.5
names(wghts) <- cols <- c("AST","CK","Daysrec","prob01")</pre>
df2 <- t(t(traindat_scaled[, cols]) * as.numeric(wghts[cols]))</pre>
traindat_wghts <- as.data.frame(df2, row.names=rownames(traindat))</pre>
colnames(traindat wghts) <- cols</pre>
###############################
# Prepare valdat.
# Compute the prob01 column.
preds01_b <- predict(g03mod, newdata=valdat)</pre>
valdat$prob01 <- as.numeric(preds01 b)</pre>
# Transform and scale valdat.
valdat$AST <- log(valdat$AST)</pre>
valdat$CK <- log(valdat$CK)</pre>
valdat$Daysrec <- sqrt(valdat$Daysrec)</pre>
valdat scaled <- scale(valdat[, -1], center=centers, scale=scales)</pre>
valdat scaled <- as.data.frame(valdat scaled, row.names=rownames(valdat))</pre>
# Apply weights to valdat.
df2 <- t(t(valdat_scaled[, cols]) * as.numeric(wghts[cols]))</pre>
valdat_wghts <- as.data.frame(df2, row.names=rownames(valdat))</pre>
colnames(valdat_wghts) <- cols</pre>
###############################
# Construct k-means model.
kmod <- suppressWarnings(kmeans(traindat_wghts, 2, iter.max = 50, nstart=</pre>
# See how the clusters are associated with Outcome.
dfout <- as.data.frame(cbind(traindat$Outcome, kmod$cluster),</pre>
                         row.names=rownames(traindat))
colnames(dfout) <- c("Outcome", "cluster")</pre>
c1_to_Outcome1 <- c1_toLevel_1(dfout)</pre>
###############################
# Apply the k-means model to valdat_wghts.
# Fook alamant of the fallowing list is a may of woldet waste
```

```
# Get the predictions for the validation set.
              cluster_assgns <- mcmapply(getCluster, valdat_asList, ctr_list,</pre>
                                             SIMPLIFY=TRUE, mc.cores=6)
              valdat_wghts$cluster <- as.numeric(cluster_assgns)</pre>
              valdat_wghts$pred_Outcome <- NA</pre>
              if(c1 to Outcome1) {
                   valdat_wghts[which(valdat_wghts$cluster==1),]$pred_Outcome <- 1</pre>
                   valdat_wghts[which(valdat_wghts$cluster==2),]$pred_Outcome <- 0</pre>
              } else {
                   valdat_wghts[which(valdat_wghts$cluster==1),]$pred_Outcome <- 0</pre>
                   valdat_wghts[which(valdat_wghts$cluster==2),]$pred_Outcome <- 1</pre>
              # Generate confusion matrix for the k-means clusters and
              # the corresponding f-score.
              preds <- as.factor(valdat_wghts$pred_Outcome)</pre>
              names(preds) <- rownames(valdat)</pre>
              ans <- get_confusion(preds, valdat[, "Outcome", drop=FALSE])</pre>
              mat <- as.matrix(ans[[1]])</pre>
              mat[is.na(mat)] <- 0</pre>
              fscore <- round(as.numeric(ans[[2]]), 4)</pre>
              acc <- round(sum(diag(mat))/floor(sum(mat)), 4)</pre>
              type2 <- round((0.4 * acc + 0.6 * ans[[2]]), 4)
              FN <- as.numeric(mat[2,1])</pre>
              FP <- as.numeric(mat[1,2])</pre>
              return(c(fscore,acc,type2,FN,FP))
In [42]: # Get scores for the hybrid model with prob01 constructed
          # from the g03 logistic model.
          set.seed(1913)
          seed_vector <- sample(1:9999, 1000, replace=FALSE)</pre>
          start <- Sys.time()</pre>
          dat_result <- get_cvInfo(seed_vector, dat)</pre>
          stop <- Sys.time()</pre>
          round(stop - start, 2)
          # Time difference of 3.53 mins
          Time difference of 3.55 mins
In [43]: dim(dat result)
          head(dat result)
          5000 - 5
          A data.frame: 6 × 5
                                       FN
                                             FP
                  fscore
                          Acc Type2
                               <dbl> <dbl>
                                           <dbl>
                  <dbl>
                         <dbl>
                                              5
           4782--1 0.4898 0.6875 0.5689
                                        20
           4782--2 0.6667 0.8000 0.7200
                                        8
                                              8
```

```
paste0("fscore mean: ", as.character(fscore_mean))
         # 0.6134
         paste0("fscore StdDev: ", as.character(fscore_sd))
         # 0.0673
         summary(dat_result$fscore)
         'fscore mean: 0.6134'
         'fscore StdDev: 0.0673'
            Min. 1st Qu. Median
                                      Mean 3rd Qu.
                                                       Max.
            0.323 0.571
                            0.615
                                     0.613
                                            0.656
                                                      0.815
In [45]: Acc_mean <- round(mean(dat_result$Acc), 4)</pre>
         Acc_sd <- round(sd(dat_result$Acc), 4)</pre>
         paste0("accuracy mean: ", as.character(Acc_mean))
         # 0.7320
         paste0("accuracy StdDev: ", as.character(Acc_sd))
         # 0.0440
         summary(dat_result$Acc)
         'accuracy mean: 0.732'
         'accuracy StdDev: 0.044'
            Min. 1st Qu. Median
                                     Mean 3rd Qu.
                                                       Max.
            0.575 0.700
                           0.738
                                     0.732 0.762
                                                      0.875
In [46]: Type2_mean <- round(mean(dat_result$Type2), 4)</pre>
         Type2_sd <- round(sd(dat_result$Type2), 4)</pre>
         paste0("Type2 mean: ", as.character(Type2_mean))
         # 0.6609
         paste0("Type2 StdDev: ", as.character(Type2_sd))
         # 0.0553
         summary(dat_result$Type2)
         'Type2 mean: 0.6609'
         'Type2 StdDev: 0.0553'
            Min. 1st Qu. Median
                                      Mean 3rd Qu.
                                                       Max.
            0.440 0.623
                            0.662
                                     0.661 0.698
                                                      0.839
```

Distribution of Type2 scores for hybrid model w/ prob01 from g03



```
In [48]: FN_mean <- round(mean(dat_result$FN), 4)
FN_sd <- round(sd(dat_result$FN), 4)
paste0("FN mean: ", as.character(FN_mean))
# 12.59
paste0("FN StdDev: ", as.character(FN_sd))
# 3.20
""
summary(dat_result$FN)</pre>
```

'FN mean: 12.589'

'FN StdDev: 3.1972'

"

```
Min. 1st Qu. Median Mean 3rd Qu. Max. 3.0 10.0 13.0 12.6 15.0 25.0
```

```
In [49]: FP_mean <- round(mean(dat_result$FP), 4)
    FP_sd <- round(sd(dat_result$FP), 4)
    paste0("FP mean: ", as.character(FP_mean))
# 8.85
    paste0("FP StdDev: ", as.character(FP_sd))
# 3.01
""
summary(dat_result$FP)</pre>
```

'FP mean: 8.8488'

'FP StdDev: 3.0077'

```
header=TRUE, row.names=1)

dim(results)

6 · 10

In [51]: # The most recent results are the last line in the table.
# The results for kmp1_g03v02 are essentially the same as
# those for kmp1_g03.
results
```

A data.frame: 6 × 10

	fscore	fscore_sd	accuracy	acc_sd	Type2	Type2_sd	FN	FN_sd	FP	FP_sd
	<dbl></dbl>									
g03	0.6183	0.0646	0.7167	0.0448	0.6577	0.0543	11.24	3.04	11.43	3.34
gbclf	0.6051	0.0665	0.7135	0.0448	0.6485	0.0552	12.03	3.23	10.89	3.29
svm02	0.6320	0.0628	0.7030	0.0472	0.6604	0.0542	14.53	3.63	9.23	3.16
km_wghts	0.6348	0.0620	0.7004	0.0462	0.6610	0.0532	8.72	2.46	15.25	3.14
kmp1_g03	0.6163	0.0676	0.7298	0.0443	0.6617	0.0556	12.22	3.06	9.40	3.00
kmp1_g03v02	0.6134	0.0673	0.7320	0.0440	0.6609	0.0553	12.59	3.20	8.85	3.01

Final Comments for the Postscript

This Postscript section shows how we can make use of tot.withinss when searching for optimal weights. approach to finding weights is more efficient than directly relying upon Type2 cross-validation scores.

However, more research needs to be done on this approach to finding weights. As we saw above, I coulc through the entire process of finding weights; it looks to be useful only for the initial grid search. The metl tot.withinss is very sensitive to how we scale the data. Without the right scaling, the grid search takes us path. Of course, before we go to far, we can run a check on the weights using Type2 (or accuracy) cross-

```
In [ ]:
```