Feature Selection Bias in Classification of High Dimensional Data

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Abstract

This vignette reproduces modified versions the calculations and the figures that appear in Section 12.3 of: Maindonald, J.H. and Braun, W.J. Data Analysis and Graphics Using R. 2^{nd} edition 2007; 3^{rd} edition 2010, Cambridge University Press. In order to reduce execution time, there are fewer folds for cross-validation and fewer simulations.

1 What groups are of interest?

The data frame golubInfo has details of the classifying factors for the 72 columns of the data set golub. The 72 observations are classified into one of the three cancer types ALL B-type (coded allB), ALL T-type (coded allT) and AML (coded aml). The two classifications that will be investigated are (1) according to tissue type and sex, given by the factor tissue.mf, and (2) according to cancer type (ALL B-type, ALL T-type, AML), given by the factor cancer.

```
library(hddplot)
Prior to version 0.53, an error in simulateScores() had
the effect that a random set of features was selected.
data(golubInfo)
with(golubInfo, table(cancer, tissue.mf))
      tissue.mf
cancer BM:NA BM:f BM:m PB:NA PB:f PB:m
                            2
  allB
           4
               19
                     10
                                 1
           0
                0
                      8
                            0
                                 0
                                       1
  allT
                2
                      3
          16
```

For the classification according to tissue type and sex (tissue.mf), restriction to the allB leukemia type and to patients whose sex is known gives a

relatively homogeneous set of data. We will define a factor tissue.mfB that classifies the allB subset of the data for which the sex of the patient is known, and for which at least two samples are available. The single allB observation that is PB:f will be omitted.

The following calculations separate out the allB subset (GolubB) of the data, and derive the factor tissue.mfB whose levels are BM:f, BM:m and PB:m:

```
attach(golubInfo)
## Identify allB samples for that are BM:f or BM:m or PB:m
subsetB <- cancer=="allB" & tissue.mf%in%c("BM:f","BM:m","PB:m")
## Form vector that identifies these as BM:f or BM:m or PB:m
tissue.mfB <- tissue.mf[subsetB, drop=TRUE]
## Separate off the relevant columns of the matrix Golub
data(Golub) # NB: variables (rows) by cases (columns)
GolubB <- Golub[,subsetB]
detach(golubInfo)</pre>
```

Distributions across features for a selection of observations

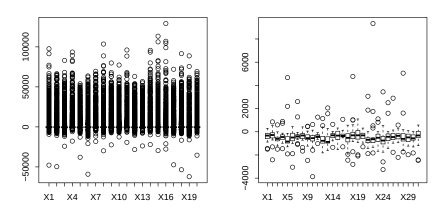


Figure 1: Coxplots of distributions across features for a selection of observations

```
## Display distributions for the first 20 observations
boxplot(data.frame(GolubB[, 1:20])) # First 20 columns (observations)
## Random selection of 20 rows (features)
boxplot(data.frame(GolubB[sample(1:7129, 20), ]))
```

2 Linear Discriminant Analysis, following variable selection

Flawed graphs

Panel A in Figure 2 shows a convincing separation between groups, based on the use of linear discriminant analysis with the 15 features that best separate the groups. Panel B, which uses random normal data to repeat the calculations for Panel A, highlights the flaws in the methodology.

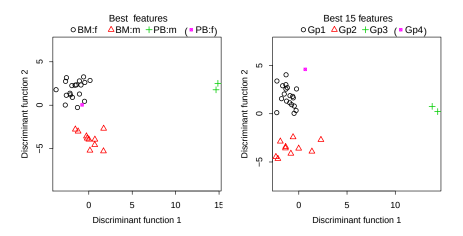


Figure 2: Panel B, with random normal data, illustrates the potential for getting spurious results with the methodology used for Panel A.

```
## Uses orderFeatures() (hddplot); see below
ord15 <- orderFeatures(GolubB, cl=tissue.mfB)[1:15]</pre>
## Panel A
dfB.ord <- data.frame(t(GolubB[ord15, ]))</pre>
## Calculations for the left panel
## Transpose to observations by features
dfB15 <- data.frame(t(GolubB[ord15, ]))</pre>
library (MASS)
dfB15.lda <- lda(dfB15, grouping=tissue.mfB)
scores <- predict(dfB15.lda, dimen=2)$x</pre>
## Scores for the single PB:f observation
df.PBf <- with(golubInfo,</pre>
  data.frame(t(Golub[ord15, tissue.mf=="PB:f" & cancer=="allB",
                      drop=FALSE])))
scores.PBf <- predict(dfB15.lda, newdata=df.PBf, dimen=2)$x</pre>
## For comparison: simulated scores
```

```
simscores <- simulateScores(nrow=7129, cl=rep(1:3, c(19,10,2)), cl.other=4, nfeatures=15, seed=41)
# Returns list elements: scores, cl, scores.other & cl.other
```

3 Distributional extremes

Calculated F-statistics (Figure 3) will be compared with the permutation distribution and with the theoretical F-distribution. Code is:

```
## In the following, B is too small for the simulation to give a
## good indication of behaviour in the extreme tail.
library(multtest, quietly=TRUE)
Loading required package: BiocGenerics
Loading required package: parallel
   Attaching package: 'BiocGenerics'
   The following objects are masked from 'package:parallel':
       clusterApply, clusterApplyLB, clusterCall, clusterEvalQ,
clusterExport, clusterMap, parApply, parCapply, parLapply,
parLapplyLB, parRapply, parSapply, parSapplyLB
   The following object is masked from 'package:stats':
       xtabs
   The following objects are masked from 'package:base':
       anyDuplicated, as.data.frame, cbind, colnames, duplicated,
eval, Filter, Find, get, intersect, lapply, Map, mapply,
match, mget, order, paste, pmax, pmax.int, pmin, pmin.int,
Position, rank, rbind, Reduce, rep.int, rownames, sapply,
setdiff, sort, table, tapply, union, unique, unlist
   Welcome to Bioconductor
       Vignettes contain introductory material; view with
'browseVignettes()'. To cite Bioconductor, see
'citation("Biobase")', and for packages
'citation("pkgname")'.
GolubB.maxT <- mt.maxT(GolubB, unclass(tissue.mfB)-1, test="f",</pre>
                       B=1000)
```

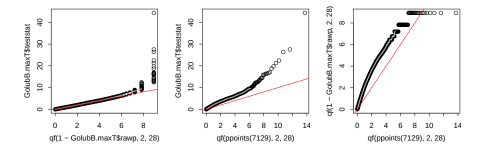


Figure 3: Compare calculated F-statistics with the permutation distribution and with the theoretical F. The theoretical F makes unrealistic normality and independence assumptions.

4 Discriminant Analysis – Training/Test

```
## Selection of features that discriminate

## ss 12.3.3: Accuracies and Scores for test data

Golub.BM <- with(golubInfo, Golub[, BM.PB=="BM"])

cancer.BM <- with(golubInfo, cancer[BM.PB=="BM"])

## Now split each of the cancer.BM categories between two subsets

## Uses divideUp(), from hddplot

gp.id <- divideUp(cancer.BM, nset=2, seed=29)

# Set seed to allow exact reproduction of the results below

table(gp.id, cancer.BM)

cancer.BM

gp.id allB allT aml

1 17 4 10

2 16 4 11
```

```
accboth <- accTrainTest(x = Golub.BM, cl = cancer.BM, traintest=gp.id, , print.progress=FALSE)

Training/test split Best accuracy, less 1SD
I (training) / II (test) 0.89 (14 features)
II (training) / I (test) 0.92 (10 features)

Training/test split Best accuracy
I (training) / II (test) 0.94 (20 features)
II (training) / II (test) 0.97 (17 features)
```

Code for plotting the figures is:

```
opar <- par(mar=c(4,4,3.1,.1))
## Use function plotTrainTest() from hddplot</pre>
```

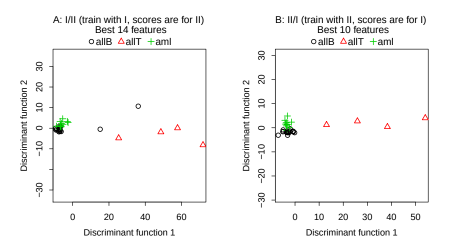


Figure 4: Panel A plots scores for the set II data, using set I for training (the I/II split), as described in the text. Panel B plots the scores for the set I data when the roles of the two sets were reversed, i.e., the split was II/I.

```
plotTrainTest(x=Golub.BM, nfeatures=c(14,10), cl=cancer.BM, traintest=gp.id)
par(opar)
```

Now compare the choice of features between I/II and II/I:

```
rbind(accboth$sub1.2[1:20],accboth$sub2.1[1:20])
     [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9]
                                                     [,10]
                                                                 [,12]
                                                           [,11]
[1,] 4050 2794 6510 6696 4342 5542 4357 5543 1207
                                                     4584
                                                            6236
                                                                  1429
[2,] 6606 4342 6510 3594 4050 6236
                                                            5542
                                                                  2061
                                    1694 1207 1268
                                                     4847
           [,14]
                 [,15]
                        [,16]
                              [,17]
                                     [,18]
                                           [,19]
                                                 [,20]
     [,13]
[1,]
      6575
            2833
                  4750
                         2335
                               1704
                                      4882
                                            6225
                                                  3544
[2,]
      5543
            4055
                  4375
                        1144
                                379
                                      6696
                                            4196
                                                   229
match(accboth$sub1.2[1:20],accboth$sub2.1[1:20])
      5 NA 3 18 2 11 NA 13 8 NA 6 NA NA NA NA NA NA NA NA NA NA
```

4.1 Cross-validation based optimum choice of features

With the number of selected features varying from 1 to 25, three different accuracy measures will be compared, for classification of the B-cell data. The plots highlight the serious bias in measures that are to an extent internal to the training data.

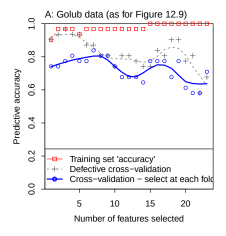
```
## Cross-validation to determine the optimum number of features
## Accuracy measure will be: tissue.mfB.cvfacc.cv
tissue.mfB.cv <- cvdisc(GolubB, cl=tissue.mfB, nfeatures=1:23,</pre>
                        nfold=c(5,1), print.progress=FALSE)
Accuracy
                   Best accuracy, less 1SD Best accuracy
(Cross-validation) 0.77 (3 features)
                                             0.84 (7 features)
  # 5-fold CV (x1)
## Defective measures will be in acc.resub (resubstitution)
## and acc.sel1 (select features prior to cross-validation)
tissue.mfB.badcv <- defectiveCVdisc(GolubB, cl=tissue.mfB,</pre>
                                    foldids=tissue.mfB.cv$folds,
                                    nfeatures=1:23, nfold=c(5,1),
                                    print.progress=FALSE)
## NB: Warning messages have been omitted
## Calculations for random normal data:
set.seed(43)
rGolubB <- matrix(rnorm(prod(dim(GolubB))), nrow=dim(GolubB)[1])</pre>
rtissue.mfB.cv <- cvdisc(rGolubB, cl=tissue.mfB, nfeatures=1:23,</pre>
                         nfold=c(5,1), print.progress=FALSE)
[1] "Input rows (features) are not named. Names"
[1] "1:7129 will be assigned."
Accuracy
                   Best accuracy, less 1SD Best accuracy
(Cross-validation) 0.46 (1 features)
                                              0.55 (1 features)
rtissue.mfB.badcv <- defectiveCVdisc(rGolubB, cl=tissue.mfB,
                                    nfeatures=1:23, nfold=c(5,1),
                                    foldids=rtissue.mfB.cv$folds,
                                    print.progress=FALSE)
[1] "Input rows (features) are not named. Names"
[1] "1:7129 will be assigned."
## This function will be used for the plots
plot.acc <- function(cv=cv1, badcv=badcv1, nseq=NULL, badnseq=NULL,</pre>
                     title="", ylab="Predictive accuracy",
```

add.legend=TRUE) {

```
maxg <- min(c(length(badcv$acc.resub), length(cv$acc.cv)))</pre>
if(is.null(nseq))nseq <- 1:maxg</pre>
plot(nseq, badcv$acc.resub[1:maxg], ylim=c(0,1), type="n",
     yaxs="i", xlab="Number of features selected", ylab=ylab)
par(xpd=T)
points(nseq, badcv$acc.resub[1:maxg], col=2, type="b", lty=2,
       pch=0, cex=0.8)
par(xpd=FALSE)
points(nseq, badcv$acc.sel1[1:maxg], col="gray40", pch=3, cex=0.8)
lines(lowess(nseq, badcv$acc.sel1[1:maxg], f=.325, iter=0),
      col="gray40", lty=2)
points(nseq, cv$acc.cv[1:maxg], col="blue", pch=1, cex=0.8)
lines(lowess(nseq, cv$acc.cv[1:maxg], f=.325, iter=0), col="blue",
xy \leftarrow par() susr[c(1,3)]
if(add.legend)
  legend(xy[1], xy[2], xjust=0, yjust=0,
         legend=c("Training set 'accuracy'",
           "Defective cross-validation",
           "Cross-validation - select at each fold"),
         lty=c(1,2,1), lwd=c(1,1,2), pch=c(0,3,1),
         col=c("red", "gray40", "blue"), cex=0.875)
mtext(side=3,line=0.35, title, adj=0)
```

Figure 5 compares three different accuracy measures, for the classification of the B-cell data. The training data measure (\square) is a severely biased measure. Cross-validation, but with features selected using all the data (+), is less severely biased. An acceptable measure of predictive accuracy (\circ) requires re-selection of features at each fold of the cross-validation. The right panel shows the performance of each of these measures when the expression values were replaced by random data.

Which features?



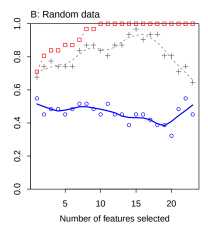


Figure 5: Comparison of different accuracy measures, in the development of a discriminant rule for the classification, into the categories BM:f, BM:m and PB:m, of the B-cell ALL data for which gender is known.

```
U29195_at 1 0 0

U91327_at 0 2 0

U49395_at 0 1 0

X00437_s_at 0 1 0

X54870_at 0 1 0

X62654_rna1_at 0 0 3

X82494_at 0 0 1
```

5 Cross-validation: bone marrow (BM) samples

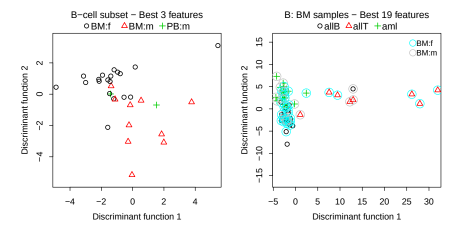


Figure 6: These plots of projections of linear discriminant analysis scores are designed to fairly reflect the performance of a linear discriminant in distinguishing between known groups in the data. The two panels relate to different subsets of the Golub data, with different groupings in the two cases. In panel B, for the classification of the 62 bone marrow (BM) samples into allB, allT, and aml, points where the sex is known are identified as male or female.

Code is:

```
prefix="B: BM samples -")
par(opar)
```