Classification of Breast Cancer Clinical Stage with Gene Expression Data

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This document presents analysis for the MAQC-II project, human breast cancer data set with boosting algorithms developed in Wang (2018a,b) and implemented in R package bst.

Dataset comes from the MicroArray Quality Control (MAQC) II project and includes 278 breast cancer samples with 164 estrogen receptor (ER) positive cases. The data files GSE20194_series_matrix.txt.gz and GSE20194_MDACC_Sample_Info.xls can be downloaded from http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?token=rhojvaiwkcsaihq&acc=GSE20194. After reading the data, some unused variables are removed. From 22283 genes, the dataset is pre-screened to obtain 3000 genes with the largest absolute values of the two-sample t-statistics. The 3000 genes are standardized.

```
# The data files below were downloaded on June 1, 2016
require("gdata")
bc <- t(read.delim("GSE20194_series_matrix.txt.gz", sep = "",</pre>
    header = FALSE, skip = 80))
colnames(bc) <- bc[1, ]</pre>
bc \leftarrow bc[-1, -c(1, 2)]
### The last column is empty with variable name
### !series_matrix_table_end, thus omitted
bc \leftarrow bc[, -22284]
mode(bc) <- "numeric" ### convert character to numeric</pre>
dat1 <- read.xls("GSE20194_MDACC_Sample_Info.xls", sheet = 1,</pre>
    header = TRUE)
y <- dat1$characteristics..ER_status
y < - ifelse(y == "P", 1, -1)
table(y)
res <- rep(NA, dim(bc)[2])
for (i in 1:dim(bc)[2]) res[i] <- abs(t.test(bc[, i] ~ y)$statistic)</pre>
### find 3000 largest absolute value of t-statistic
tmp <- order(res, decreasing = TRUE)[1:3000]
dat <- bc[, tmp]
### standardize variables
dat <- scale(dat)</pre>
```

Set up configuration parameters.

```
nrun <- 100
per \leftarrow c(0, 0.05, 0.1, 0.15)
learntype <- c("tree", "ls")[2]</pre>
tuning <- "error"</pre>
n.cores <- 4
plot.it <- TRUE
### robust tuning parameters used in bst/rbst function
s \leftarrow c(0.9, 1.01, 0.5, -0.2, 0.8, -0.5, -0.2)
nu \leftarrow c(0.01, 0.1, 0.01, rep(0.1, 4))
m <- 100 ### boosting iteration number
### whether to truncate the predicted values in each boosting
### iteration?
ctr.trun <- c(TRUE, rep(FALSE, 6))
### used in bst function
bsttype <- c("closs", "gloss", "qloss", "binom", "binom", "hinge",
    "expo")
### and corresponding labels
bsttype1 <- c("ClossBoost", "GlossBoost", "QlossBoost", "LogitBoost",
    "LogitBoost", "HingeBoost", "AdaBoost")
### used in rbst function
rbsttype <- c("closs", "gloss", "qloss", "tbinom", "binomd",
    "thinge", "texpo")
### and corresponding labels
rbsttype1 <- c("ClossBoostQM", "GlossBoostQM", "QlossBoostQM",
    "TLogitBoost", "DlogitBoost", "THingeBoost", "TAdaBoost")
```

The training data contains randomly selected 50 samples with positive estrogen receptor status and 50 samples with negative estrogen receptor status, and the rest were designated as the test data. The training data is contaminated by randomly switching response variable labels at varying pre-specified proportions per=0, 0.05, 0.1, 0.15. This process is repeated nrun=100 times. The base learner is learntype=ls (with quotes). To select optimal boosting iteration from maximum value of m=100, we run five-fold cross-validation averaging classification errors. In cross-validation, we set the number of cores for parallel computing by n.cores=4. Selected results can be plotted if plot.it=TRUE. Gradient based boosting includes ClossBoost, GlossBoost, QlossBoost, LogitBoost, HingeBoost and AdaBoost. Robust boosting using rbst contains ClossBoostQM, GlossBoostQM, QlossBoostQM, TLogitBoost, DlogitBoost, THingeBoost and TAdaBoost.

```
colnames(err.m1) <- colnames(err.m2) <- c("cont-0%", "cont-5%",</pre>
    "cont-10%", "cont-15%")
colnames(mstopcv.m1) <- colnames(mstopcv.m2) <- colnames(err.m1)</pre>
colnames(nvar.m1) <- colnames(nvar.m2) <- colnames(err.m1)</pre>
colnames(errbest.m1) <- colnames(errbest.m2) <- colnames(err.m1)</pre>
colnames(mstopbest.m1) <- colnames(mstopbest.m2) <- colnames(err.m1)</pre>
for (ii in 1:nrun) {
    set.seed(1000 + ii)
    trid <- c(sample(which(y == 1))[1:50], sample(which(y ==</pre>
        -1))[1:50])
    dtr <- dat[trid, ]</pre>
    dte <- dat[-trid, ]</pre>
    ytrold <- y[trid]
    yte <- y[-trid]</pre>
    ### number of patients/no. variables in training and test data
    dim(dtr)
    dim(dte)
    ### randomly contaminate data
    ntr <- length(trid)</pre>
    set.seed(1000 + ii)
    con <- sample(ntr)</pre>
    for (j in 1) {
        ### controls learntype i controls how many percentage of data
        ### contaminated
        for (i in 1:4) {
            ytr <- ytrold
            percon <- per[i]</pre>
            ### randomly flip labels of the samples in training set
            ### according to pre-defined contamination level
            if (percon > 0) {
               ji <- con[1:(percon * ntr)]</pre>
              ytr[ji] <- -ytrold[ji]</pre>
            dat.m1 <- bst(x = dtr, y = ytr, ctrl = bst_control(mstop = m,</pre>
               center = FALSE, trace = FALSE, nu = nu[k],
               s = s[k], trun = ctr.trun[k]), family = bsttype[k],
               learner = learntype[j])
             err1 <- predict(dat.m1, newdata = dte, newy = yte,
               type = "error")
             err1tr <- predict(dat.m1, newdata = dtr, newy = ytr,
               type = "loss")
            ### cross-validation to select best boosting iteration
             set.seed(1000 + ii)
             cvm1 \leftarrow cv.bst(x = dtr, y = ytr, K = 5, n.cores = n.cores,
               ctrl = bst_control(mstop = m, center = FALSE,
                 trace = FALSE, nu = nu[k], s = s[k], trun = ctr.trun[k]),
               family = bsttype[k], learner = learntype[j],
               main = bsttype[k], type = tuning, plot.it = FALSE)
             optmstop <- max(10, which.min(cvm1$cv))</pre>
```

```
err.m1[ii, i] <- err1[optmstop]</pre>
        nvar.m1[ii, i] <- nsel(dat.m1, optmstop)[optmstop]</pre>
        errbest.m1[ii, i] <- min(err1)</pre>
        mstopbest.m1[ii, i] <- which.min(err1)</pre>
        mstopcv.m1[ii, i] <- optmstop</pre>
        dat.m2 <- rbst(x = dtr, y = ytr, ctrl = bst_control(mstop = m,</pre>
          iter = 100, nu = nu[k], s = s[k], trun = ctr.trun[k],
          center = FALSE, trace = FALSE), rfamily = rbsttype[k],
          learner = learntype[j])
        err2 <- predict(dat.m2, newdata = dte, newy = yte,
          type = "error")
        err2tr <- predict(dat.m2, newdata = dtr, newy = ytr,
          type = "loss")
        ### cross-validation to select best boosting iteration
        set.seed(1000 + ii)
        cvm2 \leftarrow cv.rbst(x = dtr, y = ytr, K = 5, n.cores = n.cores,
          ctrl = bst_control(mstop = m, iter = 100, nu = nu[k],
            s = s[k], trun = ctr.trun[k], center = FALSE,
            trace = FALSE), rfamily = rbsttype[k], learner = learntype[j],
          main = rbsttype[k], type = tuning, plot.it = FALSE)
        optmstop <- max(10, which.min(cvm2$cv))</pre>
        err.m2[ii, i] <- err2[optmstop]</pre>
        nvar.m2[ii, i] <- nsel(dat.m2, optmstop)[optmstop]</pre>
        errbest.m2[ii, i] <- min(err2)</pre>
        mstopbest.m2[ii, i] <- which.min(err2)</pre>
        mstopcv.m2[ii, i] <- optmstop</pre>
    }
}
if (ii%%nrun == 0) {
    if (bsttype[k] %in% c("closs", "gloss", "qloss"))
        cat(paste("\nbst family ", bsttype1[k], ", s=",
          s[k], ", nu=", nu[k], sep = ""), "\n")
    if (bsttype[k] %in% c("binom", "hinge", "expo"))
        cat(paste("\nbst family ", bsttype1[k], ", nu=",
          nu[k], sep = ""), "\n")
    cat("best misclassification error from bst\n")
    print(round(apply(errbest.m1, 2, summary7), 4))
    cat("CV based misclassification error from bst\n")
    print(round(apply(err.m1, 2, summary7), 4))
    cat("best mstop with best misclassification error from bst\n")
    print(round(apply(mstopbest.m1, 2, summary7), 0))
    cat("best mstop with CV from bst\n")
    print(round(apply(mstopcv.m1, 2, summary7), 0))
    cat("nvar from bst\n")
    print(round(apply(nvar.m1, 2, summary7), 1))
    cat(paste("\nrbst family ", rbsttype1[k], ", s=",
        s[k], ", nu=", nu[k], sep = ""), "\n")
    cat("\nbest misclassification error from rbst\n")
```

```
print(round(apply(errbest.m2, 2, summary7), 4))
cat("CV based misclassification error from rbst\n")
print(round(apply(err.m2, 2, summary7), 4))
cat("best mstop with best misclassification error from rbst\n")
print(round(apply(mstopbest.m2, 2, summary7), 0))
cat("best mstop with CV from rbst\n")
print(round(apply(mstopcv.m2, 2, summary7), 0))
cat("nvar from rbst\n")
print(round(apply(nvar.m2, 2, summary7), 1))
res <- list(err.m1 = err.m1, nvar.m1 = nvar.m1, errbest.m1 = errbest.m1,
   mstopbest.m1 = mstopbest.m1, mstopcv.m1 = mstopcv.m1,
    err.m2 = err.m2, nvar.m2 = nvar.m2, errbest.m2 = errbest.m2,
   mstopbest.m2 = mstopbest.m2, mstopcv.m2 = mstopcv.m2,
    s = s[k], nu = nu[k], trun = ctr.trun[k], family = bsttype[k],
    rfamily = rbsttype[k])
if (plot.it) {
   par(mfrow = c(2, 1))
    boxplot(err.m1, main = "Misclassification error",
      subset = "", sub = bsttype1[k])
    boxplot(err.m2, main = "Misclassification error",
      subset = "", sub = rbsttype1[k])
   boxplot(nvar.m1, main = "No. variables", subset = "",
      sub = bsttype1[k])
   boxplot(nvar.m2, main = "No. variables", subset = "",
      sub = rbsttype1[k])
check <- FALSE
if (check) {
   par(mfrow = c(3, 1))
    title <- paste("percentage of contamination ",
      percon, sep = "")
    plot(err2tr, main = title, ylab = "Loss value",
     xlab = "Iteration", type = "l", lty = "dashed",
      col = "red")
    points(err1tr, type = "l", lty = "solid", col = "black")
    legend("topright", c(bsttype1[k], rbsttype1[k]),
      lty = c("solid", "dashed"), col = c("black",
    plot(err2, main = title, ylab = "Misclassification error",
     xlab = "Iteration", type = "l", lty = "dashed",
      col = "red")
    points(err1, type = "1")
    legend("bottomright", c(bsttype1[k], rbsttype1[k]),
      lty = c("solid", "dashed"), col = c("black",
        "red"))
    plot(nsel(dat.m2, m), main = title, ylab = "No. variables",
     xlab = "Iteration", lty = "dashed", col = "red",
      type = "l")
   points(nsel(dat.m1, m), ylab = "No. variables",
```

```
xlab = "Iteration", lty = "solid", type = "l",
                  col = "black")
                legend("bottomright", c(bsttype1[k], rbsttype1[k]),
                  lty = c("solid", "dashed"), col = c("black",
                    "red"))
           }
        }
    }
print(proc.time() - ptm)
sessionInfo()
## R version 3.5.2 (2018-12-20)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 18.04.1 LTS
## Matrix products: default
## BLAS: /usr/lib/x86_64-linux-gnu/blas/libblas.so.3.7.1
## LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.7.1
## locale:
  [1] LC_CTYPE=en_US.UTF-8
##
                                  LC_NUMERIC=C
  [3] LC_TIME=en_US.UTF-8
                                  LC_COLLATE=C
   [5] LC_MONETARY=en_US.UTF-8
                                  LC_MESSAGES=en_US.UTF-8
   [7] LC_PAPER=en_US.UTF-8
                                  LC_NAME=C
##
   [9] LC_ADDRESS=C
                                  LC_TELEPHONE=C
##
## [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
## attached base packages:
## [1] stats
               graphics grDevices utils
                                               datasets
## [6] methods
                base
##
## other attached packages:
## [1] knitr_1.21
## loaded via a namespace (and not attached):
## [1] compiler_3.5.2
                       magrittr_1.5
                                            formatR_1.5
## [4] tools_3.5.2
                         R.rsp_0.43.0
                                            stringi_1.2.4
## [7] R.methodsS3_1.7.1 stringr_1.3.1
                                            xfun_0.4
## [10] digest_0.6.18
                         R.cache_0.13.0
                                            R.utils_2.7.0
## [13] evaluate_0.12
                         R.oo_1.22.0
```

References

Zhu Wang. Robust boosting with truncated loss functions. *Electronic Journal of Statistics*, 12(1):599–650, 2018a. doi: 10.1214/18-EJS1404.

Zhu Wang. Quadratic majorization for nonconvex loss with applications to the boosting algorithm. Journal of Computational and Graphical Statistics, 27(3): 491-502, 2018b. doi: 10.1080/10618600.2018.1424635.