

Classification of UCI Machine Learning Datasets

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This document presents benchmark data analysis similar to [Wang \(2012\)](#) using R package **bst**. We compare the multi-class HingeBoost using three different algorithms for four benchmark data sets available from the UCI repository of machine learning data. We utilized regression trees as base learners in the HingeBoost. The number of terminal nodes is related to the depth of the tree, and the degree of interactions. To illustrate, we present the results for maximum tree depth 6.

1 Image segmentation data

```
library("bst")
```

```
tmp <- "https://archive.ics.uci.edu/ml/machine-learning-databases/image/"
dat1 <- "segmentation.data"
dat1 <- read.delim(paste(tmp, dat1, sep=""), sep=",", header=FALSE, skip=5)
dat2 <- "segmentation.test"
dat2 <- read.delim(paste(tmp, dat2, sep=""), sep=",", header=FALSE, skip=5)
dat1[,1] <- as.numeric(factor(dat1[,1]))
dat2[,1] <- as.numeric(factor(dat2[,1]))
m <- 500
dat.m1 <- mbst(x=dat1[,-1], y=dat1[,1], ctrl = bst_control(mstop=m), control.tree=list(max
err.te1 <- predict(dat.m1, newdata=dat2[,-1], newy=dat2[,1], mstop=m, type="error")
dat.m2 <- mbst(x=dat1[,-1], y=dat1[,1], ctrl = bst_control(mstop=m), control.tree=list(max
err.te2 <- predict(dat.m2, newdata=dat2[,-1], newy=dat2[,1], mstop=m, type="error")
dat.m3 <- mhingebst(x=dat1[,-1], y=dat1[,1], ctrl = bst_control(mstop=m), control.tree=lis
err.te3 <- predict(dat.m3, newdata=dat2[,-1], newy=dat2[,1], mstop=m, type="error")
plot(err.te1, type="l", xlab="Iteration", ylab="Test Error", ylim=c(0.05, 0.12))
points(err.te2, type="l", lty="dashed", col="blue")
points(err.te3, type="l", lty="dotted", col="red")
legend("topright", c("mbst_hinge", "mbst_hinge2", "mhingebst"), lty=c("solid", "dashed", "
```

2 Thyroid disease classification

```

tmp <- "http://archive.ics.uci.edu/ml/machine-learning-databases/thyroid-disease/"
dat1 <- "ann-train.data"
dat1 <- read.table(paste(tmp, dat1, sep=""))
dat2 <- "ann-test.data"
dat2 <- read.table(paste(tmp, dat2, sep=""))
m <- 400
dat.m1 <- mbst(x=dat1[,-22], y=dat1[,22], ctrl = bst_control(mstop=m), control.tree=list(m
err.te1 <- predict(dat.m1, newdata=dat2[,-22], newy=dat2[,22], mstop=m, type="error")
dat.m2 <- mbst(x=dat1[,-22], y=dat1[,22], ctrl = bst_control(mstop=m), control.tree=list(m
err.te2 <- predict(dat.m2, newdata=dat2[,-22], newy=dat2[,22], mstop=m, type="error")
dat.m3 <- mhingebst(x=dat1[,-22], y=dat1[,22], ctrl = bst_control(mstop=m), control.tree=l
err.te3 <- predict(dat.m3, newdata=dat2[,-22], newy=dat2[,22], mstop=m, type="error")
plot(err.te1, type="l", xlab="Iteration", ylab="Test Error", ylim=c(0.005, 0.01))
points(err.te2, type="l", lty="dashed", col="blue")
points(err.te3, type="l", lty="dotted", col="red")
legend("topright", c("mbst_hinge", "mbst_hinge2", "mhingebst"), lty=c("solid", "dashed", "

```

3 Satellite image classification

```

tmp <- "http://archive.ics.uci.edu/ml/machine-learning-databases/statlog/satimage/"
train <- "sat.trn"
train <- read.table(paste(tmp, train, sep=""))
test <- "sat.tst"
test <- read.table(paste(tmp, test, sep=""))
train[,37] <- as.numeric(as.factor(train[,37]))
test[,37] <- as.numeric(as.factor(test[,37]))
p <- 37
colnames(train)[1:(p-1)] <- paste("x", 1:(p-1), sep = "")
colnames(test)[1:(p-1)] <- paste("x", 1:(p-1), sep = "")
m <- 600
dat.m1 <- mbst(x=train[,-37], y=train[,37], ctrl = bst_control(mstop=m), control.tree=list
err.te1 <- predict(dat.m1, newdata=test[,-37], newy=test[,37], mstop=m, type="error")
dat.m2 <- mbst(x=train[,-37], y=train[,37], ctrl = bst_control(mstop=m), control.tree=list
err.te2 <- predict(dat.m2, newdata=test[,-37], newy=test[,37], mstop=m, type="error")
dat.m3 <- mhingebst(x=train[,-37], y=train[,37], ctrl = bst_control(mstop=m), control.tree
err.te3 <- predict(dat.m3, newdata=test[,-37], newy=test[,37], mstop=m, type="error")
plot(err.te1, type="l", xlab="Iteration", ylab="Test Error", ylim=c(0, 0.3))
points(err.te2, type="l", lty="dashed", col="blue")
points(err.te3, type="l", lty="dotted", col="red")
legend("topright", c("mbst_hinge", "mbst_hinge2", "mhingebst"), lty=c("solid", "dashed", "

```

4 Glass identification database

```

dat <- "https://archive.ics.uci.edu/ml/machine-learning-databases/glass/glass.data"
dat <- read.delim(dat, sep=",", header=FALSE)[-1]
### there is no class 4
table(dat[,10])
### must recode class label such that the class labels are consecutive, which is how the c
id <- dat[,10] > 3
dat[id, 10] <- dat[id,10] - 1
table(dat[,10])
p <- ncol(dat)
colnames(dat)[1:(p-1)] <- paste("x", 1:(p-1), sep = "")
set.seed(153)
### generate 10 balanced training data and test data, using 9 folds for training and one f
allfolds <- balanced.folds(dat[,10], nfolds=10)
omit <- allfolds[[1]]
train <- dat[-omit,]
test <- dat[omit,]
m <- 200
dat.m1 <- mbst(x=train[, -p], y=train[, p], ctrl = bst_control(mstop=m), control.tree=list(m
err.te1 <- predict(dat.m1, newdata=test[, -p], newy=test[, p], mstop=m, type="error")
dat.m2 <- mbst(x=train[, -p], y=train[, p], ctrl = bst_control(mstop=m), control.tree=list(m
err.te2 <- predict(dat.m2, newdata=test[, -p], newy=test[, p], mstop=m, type="error")
dat.m3 <- mhingebst(x=train[, -p], y=train[, p], ctrl = bst_control(mstop=m), control.tree=l
err.te3 <- predict(dat.m3, newdata=test[, -p], newy=test[, p], mstop=m, type="error")
plot(err.te1, type="l", xlab="Iteration", ylab="Test Error", ylim=c(0.15, 0.36))
points(err.te2, type="l", lty="dashed", col="blue")
points(err.te3, type="l", lty="dotted", col="red")
legend("topright", c("mbst_hinge", "mbst_hinge2", "mhingebst"), lty=c("solid", "dashed", "

```

References

Zhu Wang. Multi-class HingeBoost: Method and application to the classification of cancer types using gene expression data. *Methods of Information in Medicine*, 51(2):162–167, 2012.