STA6703 SML Take-Home Prelim, Fall 2022

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Helping functions

```
myRound <- function(x, acc=3) {mult = 10^acc; round(x*mult)/mult}</pre>
```

Problem 1

```
set.seed(0)
n = 100
m=1000

origData = rnorm(n) # case 1
z=sample(x=origData, size=n*m, replace=TRUE)
case1_mat <- matrix(z,nrow=m)

origData = rt(n,df=3); # case 2
z=sample(x=origData, size=n*m, replace=TRUE)
case2_mat <- matrix(z,nrow=m)

origData = rt(n,df=25); # case 3
z=sample(x=origData, size=n*m, replace=TRUE)
case3_mat <- matrix(z,nrow=m)</pre>
```

Problem 2

```
myCVids <- function(n, K, seed=0) {
# balanced subsets generation (subset sizes differ by at most 1)
# n is the number of observations/rows in the training set
# K is the desired number of folds (e.g., 5 or 10)
set.seed(seed);
t = floor(n/K); r = n-t*K;
id0 = rep((1:K),times=t)
ids = sample(id0,t*K)
if (r > 0) {ids = c(ids, sample(K,r))}
ids
```

```
}
# two-sample t test
column_t_test <- function(features, target){</pre>
  tests = lapply(seq(1,ncol(features)),function(x){t.test(features[,x]~target)})
  pval_lst = lapply(seq(1,length(tests)),function(x){tests[[x]]$p.value})
  return(list(tests, pval_lst))
}
# keep features with p-value <= 0.05
top_features <- function(pval_lst, p_val, p_min){</pre>
  bool_lst = lapply(seq(1,length(pval_lst)),function(x){pval_lst[[x]] < p_val})
  if(sum(as.integer(bool_lst)) <= p_min) {</pre>
    return(bool_lst)
  }else{
    pval_df = t(data.frame(pval_lst))
    row.names(pval_df) <- NULL</pre>
    bool_df=data.frame(pval_sig=matrix(unlist(bool_lst), nrow=length(bool_lst), byrow=TRUE))
    bool_df$p_val <- pval_df[,1]</pre>
    bool_df$p_top = FALSE
    sorted_res = sort(bool_df$p_val, index.return=TRUE)
    bool_df[head(sorted_res$ix,p_min),]$p_top = TRUE
    bool_df$p_select = as.logical(bool_df$pval_sig*bool_df$p_top)
    selected_df = bool_df[bool_df$p_select==TRUE,]
    feat_index_vec = as.numeric(rownames(selected_df))
  }
  return(feat_index_vec)
}
# Mis-classification ratio calculation
MCR <- function(true_vals, pred_probs, threshold=0.5){</pre>
  if(length(true_vals)!=length(pred_probs)){
    print("ERROR: predictions and true values not of same shape")
  }else{
    pred_vals = as.integer((pred_probs > threshold))
    mcr = sum(pred_vals != true_vals)/length(true_vals)
    return(mcr)
  }
}
# Generate data
set.seed(0)
n = 25
nr = n*2
Y = c(rep(1,n), rep(0,n))
p_{star_vec} = c(5,10,20,40)
```

```
i_vec = seq(5)
mcr_i_pstar_df = data.frame(matrix(0, nrow = length(p_star_vec), ncol = length(i_vec)+1))
mcr_i_pstar_df[,1] = p_star_vec
for(i in i_vec){
  nc = 200*2^5 \# nc = 6400
  M = matrix(rnorm(nr*nc),nrow=nr)
  X = M[,1:nc]
  ids = myCVids(n=nr, K=k, seed=0)
  # apply two-sample t-test
  t_test_results = column_t_test(features=X, target=Y)
  # get all features according to p-value at differing values of p_star
  mean_pstar_mcr_vec = c()
  for(p_star in p_star_vec){
    feat_index_vec = c(top_features(pval_lst=t_test_results[[2]], p_val=0.05, p_min=p_star))
    # assign(paste("feat_idx_vec_", p_star, sep = ""), feat_index_vec)
    X_pstar = X[,feat_index_vec]
    k_mcr_vec = c()
    for( k in seq(k)){
      isk = (ids == k) # k varies from 1 to K
      valid.k = which(isk)
      train.k = which(!isk)
      train_df = data.frame(Y=Y[train.k], X_pstar[train.k,])
      val_df = data.frame(Y=Y[valid.k], X_pstar[valid.k,])
      LR = glm(Y \sim .,
              data=train_df,
              family="binomial")
      LR_probs = data.frame(
                    predict(LR,
                    val_df,
                    type ="response"
                  )
      mcr = MCR(
              true_vals=val_df$Y,
              pred_probs=LR_probs[,1],
              threshold=0.5)
      k_mcr_vec = c(k_mcr_vec, mcr)
    mean_pstar_mcr = mean(k_mcr_vec)
    mean_pstar_mcr_vec = c(mean_pstar_mcr_vec, mean_pstar_mcr)
```

```
mcr_i_pstar_df[,i+1] = mean_pstar_mcr_vec

mcr_i_pstar_df = t(mcr_i_pstar_df)
rownames(mcr_i_pstar_df) = c("p*", i_vec)
knitr::kable(mcr_i_pstar_df, format = "markdown")
```

p*	5.00	10.00	20.00	40.00
1	0.14	0.08	0.08	0.34
2	0.16	0.02	0.04	0.30
3	0.16	0.02	0.04	0.30
4	0.16	0.02	0.04	0.30
5	0.16	0.02	0.04	0.30
2 3 4	0.16 0.16 0.16	0.02 0.02 0.02	0.04 0.04 0.04	0.3 0.3 0.3

Problem 3

Problem 4

```
genData <- function(n, seed=0) {
set.seed(seed)
x = seq(-1,1,length.out=n)
y = x - x^2 + 2*rnorm(n) # true sigma = 2;
out.df = data.frame(x=x, y=y)
out.df
}
train.df = genData(n=200,seed=100)
test.df = genData(400)</pre>
```

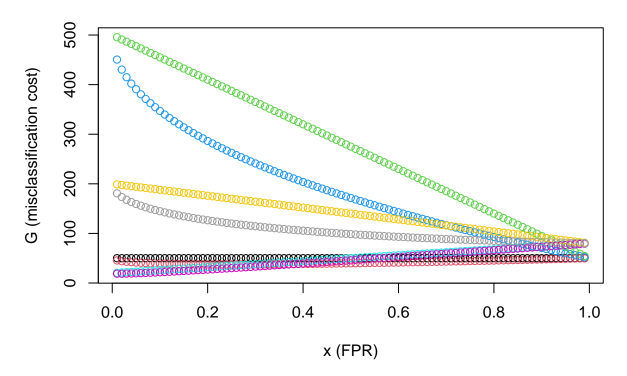
Problem 5

```
# assume values for x and cfp
# x = 0.25
cfp=1
n=100

x_vec=c()
A_vec=c()
C_vec=c()
R_vec=c()
FPR_vec=c()
TPR_vec=c()
```

```
G_vec=c()
for (x in seq(0.01,0.99,0.01)) {
  A=c(0.5, 0.2)
  C=c(cfp, 10*cfp)
  R=c(x, sqrt(x))
  Ai=0
  for(q in A){
   Ai = Ai + 1
   P=n*q
   N=n*(1-q)
    Ri=0
    for(tpr in R){
      Ri=Ri+1
      # calculate TP, FP, TN, and FN with regards to x
      TP = tpr*P
      FN = P-TP
      FP = x*N
      TN = N-FP
      FPR = x
      TPR = tpr
      Ci=0
      for(cfn in C){
        Ci=Ci+1
        G = cfn*FN + cfp*FP
        x_{\text{vec=c}}(x_{\text{vec}},x)
        A_vec=c(A_vec,Ai)
        C_vec=c(C_vec,Ci)
        R_vec=c(R_vec,Ri)
        FPR_vec=c(FPR_vec,FPR)
        TPR_vec=c(TPR_vec,TPR)
        G_vec=c(G_vec,G)
        # print(paste("(A:",as.character(Ai),")"))
        # print(paste("(C:",as.character(Ci),")"))
        # print(paste("(R:",as.character(Ri),")"))
        # print(paste("FPR = ", FPR))
        # print(paste("TPR = ", TPR))
        # print(paste("G = ", G))
        # print("----")
     }
  }
```

```
}
results_df = data.frame(x_vec,
                           A_vec,
                           C_vec,
                           R_vec,
                           FPR_vec,
                           TPR_vec,
                           G_vec)
results_df$design_id <- paste(results_df$A_vec,
                               results_df$C_vec,
                               results_df$R_vec)
# Objective function score visualization
plot(x=results_df$x_vec,
     y=results_df$G_vec,
     col=factor(results_df$design_id),
     ylab="G (misclassification cost)",
     xlab="x (FPR)")
```

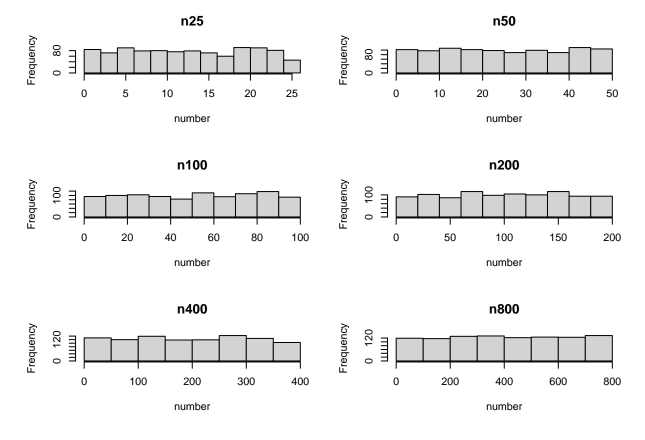


The best classifier is the one that minimizes the objective function the best over values of x. In this case it was one with the design combinations of (A:1 C1: R:2) or (A:2 C1: R:2). For higher values of x an unbalanced population gives higher values from the objective function. A1 is thus better with even populations when. This is because with an uneven class ratio the mis-classification rate increases for higher values of x because a higher x value means a higher false positive count. Even when the false

negative count is low. THis classifier is therefore not stable for all values of x and we choose the (A:1 C1: R:2) combination over the (A:2 C1: R:2) combination. Even though the (A:2 C1: R:2) combination clearly performs better at lower values of x. This in turn increased the mis-classifications calculated in the objective function. All classifiers that had a cfn that was 10 times the cfp resulted in an objective function that was orders of magnitude larger than the other classifiers. This makes intuitive senses as it dramatically increases the net cost of mis-classifications. With a TPR that is square rooted the objective function consistently returns a lower mis-classifications score. R2 is thus better. This is because, when the square root of the TPR is used to derive True positives they are higher than than when the TPR is not square rooted. This in turn decreases the amount of mis-classifications.

Problem 6

```
# Generate data
set.seed(0)
n_{\text{vec}} = c(25, 50, 100, 200, 400, 800)
m = 1000
df = data.frame(m=1:m)
table_df = data.frame(n=n_vec) # results table
mean uniq vec = c()
sd_uniq_vec = c()
for (n in n vec){
  I = seq(n)
  samp_vec = c(sample(x=I, size=m ,replace=TRUE))
  df[paste("n",n , sep="")] = samp_vec
  mean_uniq_vec = c(mean_uniq_vec, mean(unique(samp_vec)))
  sd_uniq_vec = c(sd_uniq_vec, sd(unique(samp_vec)))
}
#visualize data
par(mfrow=c(3,2))
for(i in names(df)[2:7]){
  hist(df[[i]],
       xlab = "number",
       main=i)
}
```



make table of results
<pre>table_df\$Mean_Unique = myRound(mean_uniq_vec, acc=2)</pre>
<pre>table_df\$SD_Unique = myRound(sd_uniq_vec, acc=2) table_df = data.frame(t(table_df))</pre>
<pre>knitr::kable(table_df, format = "markdown")</pre>

	X1	X2	Х3	X4	X5	X6
n			100.00			
Mean_Unique	13.00	25.50	50.50	100.75	200.02	408.84
SD_Unique	7.36	14.58	29.01	57.92	116.29	230.78