```
opts_chunk$set (warning = FALSE, message = FALSE, tidy=TRUE, echo=TRUE)
options(warn = -1)
rm(list=ls())
library(survival)
library(knitr)
library(kableExtra)
library(glmnet)
## Loading required package: Matrix
## Loaded glmnet 4.1-7
library(ggplot2)
# Following loaded in "forest_search_v0.R"
suppressMessages(library(randomForest))
#library(SPlit)
library(grf)
library(policytree)
library(DiagrammeR)
#library(cowplot)
library(data.table)
library(plyr)
library(aVirtualTwins)
# Not sure formatR is needed?
#library(formatR)
suppressMessages(library(gridExtra))
library(speff2trial)
## Loading required package: leaps
# Location where code is stored
# Modified for MAC
codepath<-c("/Users/larryleon/Documents/GitHub/forestSearch//R/")</pre>
source(paste0(codepath, "source_forestsearch_v0.R"))
source_fs_functions(file_loc=codepath)
```

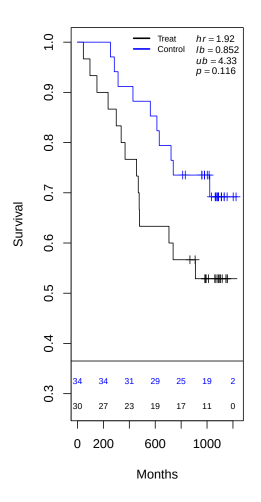
```
t.start.all <- proc.time()[3]
# GRF analysis To guide selection of binary cutpoints
df.analysis <- subset(ACTG175, arms %in% c(2, 3))

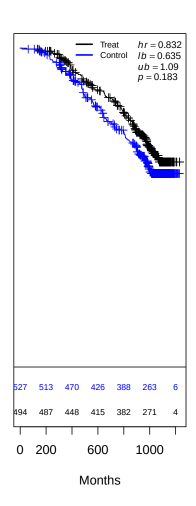
df.analysis <- within(df.analysis, {
   id <- as.numeric(c(1:nrow(df.analysis)))
   time_days <- days
   treat <- ifelse(arms == 2, 1, 0)
})

# plot(survfit(Surv(time_days,cens)~treat,data=df.analysis))
coxph(Surv(time_days, cens)~ treat, data = df.analysis)</pre>
```

```
## Call:
## coxph(formula = Surv(time_days, cens) ~ treat, data = df.analysis)
##
           coef exp(coef) se(coef)
##
## Likelihood ratio test=0.72 on 1 df, p=0.3974
## n= 1085, number of events= 237
confounders.name <- c("age", "wtkg", "karnof", "cd40", "cd80", "hemo", "homo", "drugs",
   "race", "gender", "oprior", "symptom")
outcome.name <- c("time_days")</pre>
event.name <- c("cens")
id.name <- c("id")</pre>
treat.name <- c("treat")</pre>
n.min <- 60
dmin.grf <- 12
frac.tau <- 0.8
grf.est <- grf.subg.harm.survival(data = df.analysis, confounders.name = confounders.name,
   outcome.name = outcome.name, event.name = event.name, id.name = id.name, treat.name = treat.name,
   n.min = n.min, dmin.grf = dmin.grf, frac.tau = frac.tau, details = TRUE)
## tau= 816.8
##
     leaf.node control.mean control.size control.se treated.mean treated.size
## 1
             2 -22.361327
                           827.000000
                                        9.320253
                                                  22.361327
                                                                827.000000
## 2
             3
                21.962213
                             258.000000 20.665085
                                                   -21.962213
                                                                258.000000
## 3
            4
               17.772840
                           172.000000 18.488992
                                                  -17.772840
                                                               172.000000
## 4
            5
               -81.211528
                           126.000000 28.148943
                                                   81.211528
                                                               126.000000
            6
                -30.418740
                            518.000000 11.170055
                                                    30.418740
## 5
                                                                518.000000
            7
## 6
                37.569013
                           269.000000 20.568764 -37.569013
                                                                269.000000
           10 -40.208330 456.000000 12.993950 40.208330
## 31
                                                               456.000000
                                                    -8.826842
## 41
                 8.826842
                           181.000000 10.468789
           11
                                                               181.000000
           12
                -82.358293
                                                    82.358293
## 51
                            112.000000 29.296319
                                                                112.000000
## 61
           13 74.238683 124.000000 26.034823 -74.238683 124.000000
## 7
           14 105.243703
                           64.000000 56.116526 -105.243703
                                                               64.000000
## 8
           15 -72.038631
                             81.000000 30.510745
                                                  72.038631
                                                                 81.000000
##
      treated.se
                      diff depth
## 1
       9.320253 -44.72265
## 2
       20.665085 43.92443
                               1
                 35.54568
## 3
       18.488992
## 4
       28.148943 -162.42306
                               2
## 5
       11.170055 -60.83748
       20.568764
                 75.13803
                               2
## 6
## 31
       12.993950 -80.41666
                               3
## 41
       10.468789
                 17.65368
                               3
## 51
       29.296319 -164.71659
       26.034823 148.47737
## 61
                               3
## 7
       56.116526 210.48741
## 8
       30.510745 -144.07726
                               3
   leaf.node control.mean control.size control.se treated.mean treated.size
                              64.00000 56.11653 -105.24370
## 7
                105.24370
                                                                 64.00000
         14
   treated.se
                  diff depth
## 7 56.11653 210.4874
cat("Truncation point for RMST:", c(grf.est$tau.rmst), "\n")
```

```
## Truncation point for RMST: 816.8
# Plot manually
# plot(grf.est£tree)
# plot(qrf.est£tree1)
# plot(grf.est£tree2)
# plot(grf.est£tree3)
df0.grf <- subset(grf.est$data, treat.recommend == 0)</pre>
df1.grf <- subset(grf.est$data, treat.recommend == 1)</pre>
# Terminal leaf corresponding to selected SG
cat("Terminal leaf:", c(grf.est$sg.harm.id), "\n")
## Terminal leaf: karnof <= 90
# action=1 --> recommend control
# Manually identify the subgroup looking at tree and terminal leaf
print(dim(df0.grf))
## [1] 64 34
check <- subset(df.analysis, karnof <= 90 & cd80 > 1034 & age > 37)
print(dim(check))
## [1] 64 29
# plot(survfit(Surv(time_days,cens)~treat,data=df.analysis))
# coxph(Surv(time_days,cens)~treat,data=df.analysis)
save(grf.est, file = "output/grf_actg_Arms_2vs3_final.Rdata")
## Error in gzfile(file, "wb"): cannot open the connection
t.done <- proc.time()[3]</pre>
t.min \leftarrow (t.done - t.start.all)/60
cat("Minutes and hours for GRF estimation", c(t.min, t.min/60), "\n")
## Minutes and hours for GRF estimation 2.924717 0.04874528
t.start <- proc.time()[3]</pre>
cat("GRF variables in selected tree", "\n")
## GRF variables in selected tree
print(grf.est$tree.names)
## [1] "age"
                "cd80"
                         "wtkg"
                                 "cd40"
                                            "karnof"
cat("GRF cuts wrt selected tree:", "\n")
```



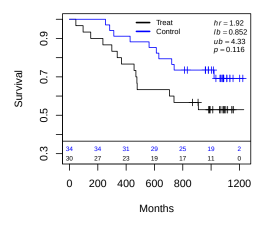


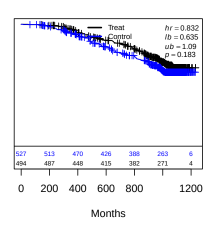
```
conflasso.name <- confounders.name[which(m$beta != 0)]</pre>
cat("Cox-LASSO selected:", c(conflasso.name), "\n")
## Cox-LASSO selected: age wtkg karnof cd40 cd80 drugs symptom
cat("GRF cuts wrt selected tree:", "\n")
## GRF cuts wrt selected tree:
print(grf.est$tree.cuts)
## [1] "age <= 37"
                        "cd80 <= 499" "cd80 <= 1034" "wtkg <= 64.64"
## [5] "cd40 <= 417" "cd80 <= 680" "karnof <= 90"
# Considering continuous factors per GRF cuts Only also considering drugs and
# symptom per lasso
df.analysis <- within(df.analysis, {</pre>
    z1a <- ifelse(age <= 37, 1, 0)
    z1b <- ifelse(age <= median(age), 1, 0)</pre>
    z2 <- ifelse(wtkg <= 65, 1, 0)
    z3 <- ifelse(karnof <= 90, 1, 0)
    z4 <- ifelse(cd40 <= 417, 1, 0)
    z5a <- ifelse(cd80 <= 499, 1, 0)
    z5b <- ifelse(cd80 <= 680, 1, 0)
    z5c <- ifelse(cd80 <= 1034, 1, 0)
   # z6<-hemo z7<-homo
    z8 <- drugs
    # z9<-race z10<-gender z11<-oprior
    z12 <- symptom
   # Convert to factors
    v1a <- as.factor(z1a)</pre>
    v1b <- as.factor(z1b)</pre>
   v2 <- as.factor(z2)</pre>
   v3 <- as.factor(z3)
    v4 <- as.factor(z4)
   v5a <- as.factor(z5a)
    v5b <- as.factor(z5b)
   v5c <- as.factor(z5c)
    v6 <- as.factor(z8)
    v7 <- as.factor(z12)
})
FSconfounders.name <- c("v1a", "v1b", "v2", "v3", "v4", "v5a", "v5b", "v5c", "v6",
    "v7")
outcome.name <- c("time_days")</pre>
event.name <- c("cens")</pre>
id.name <- c("id")</pre>
treat.name <- c("treat")</pre>
df.confounders <- df.analysis[, FSconfounders.name]</pre>
df.confounders <- dummy(df.confounders)</pre>
hr.threshold <- 1.5 # Initital candidates</pre>
hr.consistency <- 1.25 # Candidates for many splits
```

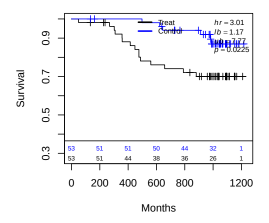
```
pconsistency.threshold <- 0.9
maxk <- 4
# maxk is max # of covariates in combination Since we want to allow generation
# of intervals for single covariate allowing for 4 can yield v1, v2 (say), and
# v3,v4 with v3 and v4 generating intervals for a single covariate
# Limit timing for forestsearch
max.minutes <- 60
nmin.fs <- 60
# stop.threshold<-0.60 # If any sq meets this, then choose this (stop here);
m1.threshold <- Inf # Turning this off (Default)</pre>
stop.threshold <- 1
# =1 will run through all sg's meeting HR criteria
fs.splits <- 1000 # How many times to split for consistency
# vi is % factor is selected in cross-validation --> higher more important
vi.grf.min <- 0.2</pre>
# Null, turns off grf screening Set to 5 for this heavily censored data
d.min <- 5 # Min number of events for both arms (d0.min=d1.min=d.min)
# default=5
sg_focus <- "hr"
split_method <- "Random"</pre>
pstop_futile <- 0.3</pre>
{\it \# Stops \ the \ consistency \ evaluation \ after \ first \ subgroup \ with \ consistency \ below}
# pstop_futile With idea that since SG's are sorted by hazard ratio estimates,
# once consistency is below pstop_futile it seems unlikely that SG's with lower
# hr's will reach the required consistency criterion
fs.est <- forestsearch(df = df.analysis, confounders.name = FSconfounders.name, df.predict = df.analysi
   details = TRUE, sg_focus = sg_focus, split_method = split_method, pstop_futile = pstop_futile,
   outcome.name = outcome.name, treat.name = treat.name, event.name = event.name,
    id.name = id.name, n.min = nmin.fs, hr.threshold = hr.threshold, hr.consistency = hr.consistency,
   fs.splits = fs.splits, stop.threshold = stop.threshold, d0.min = d.min, d1.min = d.min,
   pconsistency.threshold = pconsistency.threshold, max.minutes = max.minutes, maxk = maxk,
   plot.sg = FALSE, vi.grf.min = vi.grf.min)
## Confounders per grf screening v2 v7 v1a v3 v5b v6 v5c v1b v4 v5a
## Number of possible subgroups= 1048575
## Number of possible subgroups (in millions)= 1.048575
## # of subgroups based on # variables > k.max and excluded 1042380
## k.max = 4
## Events criteria for control, exp= 5 5
## # of subgroups with events less than criteria: control, experimental 3634 3920
## # of subgroups meeting all criteria = 1429
## # of subgroups fitted (Cox model estimable) = 1429
## Minutes= 0.1171833
## Number of criteria not met for subgroup evaluation
## crit.failure
                         2
                                 3
                1
        0
              272
## 1043809
                      3246
                               500
                                       748
## Number of subgroups meeting HR threshold 89
## Subgroups (1st 10) meeting overall screening thresholds (HR, m1) sorted by focus: (m1,sg_focus)= Inf
      K n E d1 m1 m0 HR L(HR) U(HR) v2.0 v2.1 v7.0 v7.1 v1a.0 v1a.1 v3.0
## 1: 4 106 21 15 Inf Inf 3.01 1.17 7.77 0 0 0 0 1 0 0
```

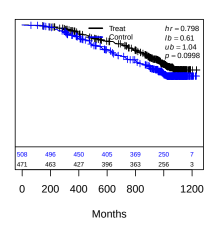
```
2: 4 99 16 11 Inf Inf 2.72 0.95 7.84
    3: 3 124 23 16 Inf Inf 2.62
                                   1.08
                                         6.36
                                                  0
                                                        0
                                                             0
                                                                   0
                                                                                     0
    4: 4 88 30 20 Inf Inf 2.49
                                                                   0
                                   1.17
                                         5.34
                                                  0
                                                        0
                                                             0
                                                                         1
                                                                                     0
    5: 4 97 29 18 Inf Inf 2.22
                                   1.05
                                         4.70
                                                  0
                                                             0
    6: 4 110 19 13 Inf Inf 2.20
                                   0.84
                                         5.80
                                                  0
                                                             1
                                                                  0
                                                                                     1
                                                                         1
    7: 4 98 31 19 Inf Inf 2.05
                                   0.99
                                         4.22
                                                  1
                                                        0
                                                             0
                                                                   0
                                                                         1
                                                                                     0
    8: 4 146 33 18 Inf Inf 2.02
                                   1.02
                                         4.01
                                                  1
                                                        0
                                                             0
                                                                   \cap
                                                                               \cap
    9: 3 120 36 22 Inf Inf 2.02 1.03 3.94
                                                  0
                                                        \cap
                                                             0
                                                                                     0
## 10: 4 157 33 22 Inf Inf 2.01 0.97 4.14
                                                  0
                                                        0
                                                             1
                                                                   0
                                                                         1
       v3.1 v5b.0 v5b.1 v6.0 v6.1 v5c.0 v5c.1 v1b.0 v1b.1 v4.0 v4.1 v5a.0 v5a.1
##
##
          0
                 1
                       0
                            1
                                  0
                                        0
                                               1
                                                      0
                                                            0
                                                                  0
                                                                       0
    1:
##
    2:
                       0
                                  0
                                         0
                                               1
                                                      0
##
   3:
          0
                 1
                       0
                             0
                                  0
                                         0
                                                      0
                                                            0
                                                                  0
                                                                             0
                                                                                    \cap
                                               1
                                                                       0
##
    4:
          1
                 1
                       0
                             0
                                  0
                                         0
                                               0
                                                      0
                                                            0
                                                                  0
                                                                             0
                                                                                    0
##
    5:
                                               0
                                                     0
          1
                 1
                       \cap
                                  0
                                         0
                                                            \cap
                                                                 \cap
                                                                       \cap
                                                                             0
                                                                                    0
                             1
##
   6:
                       0
                                  0
                                         0
                                                     0
                                                                 0
   7:
##
          1
                 1
                       0
                             0
                                  0
                                         0
                                               0
                                                     0
                                                            0
                                                                 0
                                                                       0
                                                                             0
                                                                                    0
##
    8:
          1
                 0
                       0
                                  0
                                         0
                                               0
                                                     0
                                                            1
                                                                 0
                             1
##
  9:
          1
                 1
                       0
                             \cap
                                  0
                                         \cap
                                               0
                                                     0
                                                            \cap
                                                                 0
                                                                       \cap
                                                                             \cap
                                                                                    0
                                                     0
## 10:
## Consistency 0.95
## Splitting method, # of splits= Random 1000
## Model, % Consistency Met= v1a.0 v5b.0 v6.0 v5c.1 0.95
## Consistency 0.631
## Consistency 0.913
## Splitting method, # of splits= Random 1000
## Model, % Consistency Met= v1a.0 v5b.0 v5c.1 0.913
## Consistency 0.941
## Splitting method, # of splits= Random 1000
## Model, % Consistency Met= v1a.0 v3.1 v5b.0 v4.1 0.941
## Consistency 0.641
## Consistency 0.333
## Consistency 0.533
## Consistency 0.542
## Consistency 0.493
## Consistency 0.404
## Consistency 0.501
## Consistency 0.835
## Consistency 0.835
## Consistency 0.17
## Number of subgroups meeting consistency criteria= 3
      p.consistency Nsg group.id m.index K M.1
                                                      M.2
                                                             М.З
## 1:
               0.95 106
                                21
                                         1 4 v1a.0 v5b.0 v6.0 v5c.1
## 2:
               0.913 124
                                23
                                         3 3 v1a.0 v5b.0 v5c.1
## 3:
               0.941 88
                                76
                                         4 4 v1a.0 v3.1 v5b.0
                                                                 v4.1
      p.consistency Nsg group.id m.index K
                                                M.1
                                                      M.2
                                21
## 1:
               0.950 106
                                         1 4 v1a.0 v5b.0 v6.0 v5c.1
## 2:
               0.941 88
                                76
                                         4 4 v1a.0 v3.1 v5b.0 v4.1
## 3:
               0.913 124
                                23
                                         3 3 v1a.0 v5b.0 v5c.1
xx <- fs.est$find.grps$out.found$hr.subgroups</pre>
covs.found \leftarrow xx[, -c(1:10)]
covs.most <- apply(covs.found, 2, sum)</pre>
covs.most <- covs.most[covs.most > 0]
print(covs.most)
```

```
## v2.0 v7.0 v7.1 v1a.0 v1a.1 v3.0 v3.1 v5b.0 v5b.1 v6.0 v5c.0 v5c.1 v1b.0
   36 17 1 22 16
                                  7 48 23 17
                                                        30 19 15
## v1b.1 v4.0 v4.1 v5a.0 v5a.1
   33 9 16 10
print(fs.est$grp.consistency$result)
     p.consistency Nsg group.id m.index K M.1 M.2 M.3 M.4
           0.950 106 21 1 4 v1a.0 v5b.0 v6.0 v5c.1
## 1:
## 2:
             0.941 88
                             76
                                      4 4 v1a.0 v3.1 v5b.0 v4.1
## 3:
                            23
                                      3 3 v1a.0 v5b.0 v5c.1
             0.913 124
df0.fs <- subset(fs.est$df.pred, treat.recommend == 0)
df1.fs <- subset(fs.est$df.pred, treat.recommend == 1)</pre>
save(fs.est, df.analysis, FSconfounders.name, file = "output/fs_actg_Arms_2vs3_final.Rdata")
## Error in gzfile(file, "wb"): cannot open the connection
t.done <- proc.time()[3]</pre>
t.min \leftarrow (t.done - t.start)/60
cat("Minutes and hours for FS estimation", c(t.min, t.min/60), "\n")
## Minutes and hours for FS estimation 0.4067667 0.006779444
t.start <- proc.time()[3]</pre>
# Note, the elements above will need to be re-initiated if running separate
# from above E.g., outcome.names, event.name, ... hr.threshold, etc.
# load('output/fs_actq_Arms_2vs3_final.Rdata')
library(doParallel)
registerDoParallel(parallel::detectCores(logical = FALSE))
cox.formula.boot <- as.formula(paste("Surv(time_days,cens)~treat"))</pre>
split_method <- "Random"</pre>
est.loghr <- TRUE
confounders.name <- FSconfounders.name</pre>
stop.threshold <- 0.99
fs.splits <- 1000
max.minutes <- 6
# Suggest running 50, first ... to get timing estimate
NB <- 2000
df_temp <- fs.est$df.pred[, c("id", "treat.recommend")]</pre>
dfa <- merge(df.analysis, df_temp, by = "id")</pre>
df_boot_analysis <- dfa</pre>
fitH <- get_Cox_sg(df_sg = subset(df_boot_analysis, treat.recommend == 0), cox.formula = cox.formula.bo
    est.loghr = est.loghr)
H_obs <- fitH$est_obs # log(hr) scale
seH_obs <- fitH$se_obs
```









if (dim(Ystar_mat)[1] != NB | dim(Ystar_mat)[2] != nrow(df_boot_analysis)) stop("Dimension of Ystar_mat

```
tB.start <- proc.time()[3]</pre>
# Bootstraps
resB <- bootPar({</pre>
   ans <- fsboot_forparallel(boot)</pre>
}, boots = NB, seed = 8316951, counter = "boot", export = fun_arg_list_boot)
tB.now <- proc.time()[3]
tB.min <- (tB.now - tB.start)/60
doParallel::stopImplicitCluster()
cat("Minutes for Boots", c(NB, tB.min), "\n")
## Minutes for Boots 2000 83.4098
cat("Projection per 100", c(tB.min * (100/NB)), "\n")
## Projection per 100 4.17049
cat("Propn bootstrap subgroups found =", c(sum(!is.na(resB$H_biasadj_1))/NB), "\n")
## Propn bootstrap subgroups found = 0.981
# How many timmed out
cat("Number timmed out=", c(sum(is.na(resB$H_biasadj_1) & resB$tmins_search > max.minutes)),
   "\n"
## Number timmed out= 0
H_estimates <- get_dfRes(Hobs = H_obs, seHobs = seH_obs, H1_adj = resB$H_biasadj_1,</pre>
   ystar = Ystar_mat, cov_method = "standard", cov_trim = 0.05)
Hc_estimates <- get_dfRes(Hobs = Hc_obs, seHobs = seHc_obs, H1_adj = resB$Hc_biasadj_1,</pre>
   ystar = Ystar_mat, cov_method = "standard", cov_trim = 0.05)
print(H_estimates)
                sdHO HO_lower HO_upper
                                            H1
                                                     sdH1 H1_lower H1_upper
print(Hc_estimates)
            HO sdHO HO_lower HO_upper H1
                                                     sdH1 H1_lower H1_upper
## 1: 0.7977977 0.1095 0.6096253 1.044053 0.86181 0.1204216 0.6553483 1.133315
save(fs.est, Ystar_mat, resB, H_estimates, Hc_estimates, df_boot_analysis, file = "output/fsBoot_actg_A
## Error in gzfile(file, "wb"): cannot open the connection
t.done <- proc.time()[3]</pre>
t.min \leftarrow (t.done - t.start)/60
```

cat("Minutes and hours for FS bootstrap", c(t.min, t.min/60), "\n")

Minutes and hours for FS bootstrap 83.43738 1.390623

```
df0.fs <- subset(fs.est$df.pred, treat.recommend == 0)</pre>
df1.fs <- subset(fs.est$df.pred, treat.recommend == 1)</pre>
# ITT analysis
cox_itt <- summary(coxph(Surv(time_days, cens) ~ treat, data = fs.est$df.pred))$conf.int</pre>
# ITT estimates
resITT <- c(round(cox_itt[c(1, 3, 4)], 2), nrow(fs.est$df.pred))</pre>
# Forest Search Un-adjusted
Hstat <- c(unlist(H_estimates))[c(1, 3, 4)]</pre>
resH_obs <- c(c(Hstat), nrow(df0.fs))</pre>
# Bias-corrected
Hstat <- c(unlist(H_estimates))[c(5, 7, 8)]</pre>
resH_bc <- c(c(Hstat), nrow(df0.fs))</pre>
Hstat2 <- c(unlist(H_estimates))[c(5, 7, 8)]</pre>
Hstat2 <- round(Hstat2, 2)</pre>
a <- paste0(Hstat2[1], " [")</pre>
a <- paste0(a, Hstat2[2])
a <- paste0(a, ",")
a <- paste0(a, Hstat2[3])
a <- paste0(a, "]")
H_bc2 \leftarrow c(a)
# Un-adjusted
Hcstat \leftarrow c(unlist(Hc_estimates))[c(1, 3, 4)]
resHc_obs <- c(c(Hcstat), nrow(df1.fs))</pre>
# Bias-corrected
Hcstat <- c(unlist(Hc_estimates))[c(5, 7, 8)]</pre>
resHc_bc <- c(c(Hcstat), nrow(df1.fs))</pre>
Hcstat2 \leftarrow c(unlist(Hc_estimates))[c(5, 7, 8)]
Hcstat2 <- round(Hcstat2, 2)</pre>
a <- paste0(Hcstat2[1], " [")</pre>
a <- paste0(a, Hcstat2[2])</pre>
a <- paste0(a, ",")
a <- paste0(a, Hcstat2[3])
a <- paste0(a, "]")
Hc_bc2 \leftarrow c(a)
res <- rbind(resITT, resH_obs, resH_bc, resHc_obs, resHc_bc)</pre>
resf <- as.data.frame(res)</pre>
colnames(resf) <- c("HR Estimate", "Lower", "Upper", "$\\#$ Subjects")</pre>
rnH \leftarrow c("\$\\ hat{H}$", "$\\ hat{H}_{bc}$")
rnHc \leftarrow c("\$\\ hat{H}^{c}$", "$\\ hat{H}^{c}_{bc}$")
rnItt <- c("ITT")</pre>
rownames(resf) <- c(rnItt, rnH, rnHc)</pre>
```

Table 1: ACTG-175 FS Analysis: Cox hazard ratio (HR) estimates for the ITT population and subgroups H and H^c . Cox model estimates are based on subgroups: H true (knowing the actual subgroup, a-priori); the estimated subgroup \hat{H} ; and the bootstrap (B=2,000) bias-correction to \hat{H} estimates, denoted \hat{H}_{bc} . Estimates for the complement H^c are defined analogously. The number of subjects in each population (# Subjects) are listed.

	HR Estimate	Lower	Upper	# Subjects
ITT				
ITT	0.900	0.690	1.160	1085
H subgroup estimates				
\hat{H}	3.012	1.168	7.769	106
\hat{H}_{bc}	1.535	1.054	2.238	106
H-complement subgroup estimates				
\hat{H}^c	0.798	0.610	1.044	979
\hat{H}^c_{bc}	0.862	0.655	1.133	979

```
t.done <- proc.time()[3]
t.min <- (t.done - t.start.all)/60
cat("Minutes and hours to finish", c(t.min, t.min/60), "\n")
## Minutes and hours to finish 86.77752 1.446292
cat("Machine=", c(Sys.info()[[4]]), "\n")
## Machine= Mac-Studio-3.local
cat("Number of cores=", c(detectCores(logical = FALSE)), "\n")
## Number of cores= 10</pre>
```