Build Survival Model: Random Survival Forest

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<pre>library(tidyverse) library(survival) library(randomForestSRC) library(caret) library(survcomp) library(parallel) library(doParallel) library(mcprogress) # wrap mclapply with progress bar. library(kableExtra) # include knitr automatically source("/work/users/y/u/yuukias/BIOS-Material/BIOS992/utils/csv_utils.r") # * Don't use setwd() for Quarto documents! # setwd("/work/users/y/u/yuukias/BIOS-Material/BIOS992/data")</pre>	
<pre>adjust_type <- ifelse(exists("params"), params\$adjust_type, "minimal") #</pre>	

```
# string of parameters
adjust_type_str <- switch(adjust_type,
    minimal = "minimal",
    partial = "partial",
    full = "full"
)
print(paste0("Model Adjustment Type: ", adjust_type_str))</pre>
```

[1] "Model Adjustment Type: minimal"

```
impute_type_str <- switch(impute_type,
          unimputed = "unimputed",
          imputed = "imputed"
)
print(paste0("Data Imputation Type: ", impute_type_str))</pre>
```

[1] "Data Imputation Type: unimputed"

Load Data

```
if (include_statin == "yes") {
    data_train <-
    read.csv(paste0("/work/users/y/u/yuukias/BIOS-Material/BIOS992/data/train_data_",
    impute_type_str, "_statin.csv"),
        header = TRUE
    )
} else {
    data_train <-
    read.csv(paste0("/work/users/y/u/yuukias/BIOS-Material/BIOS992/data/train_data_",
    impute_type_str, ".csv"),
        header = TRUE
    )
}
data_train <- data_train[, -1] # the first column is the index generated by
    sklearn
(dim(data_train))</pre>
```

[1] 28127 100

```
data <- select_subset(data_train, type = adjust_type)
(dim(data))</pre>
```

[1] 28127 48

colnames(data)

```
[1] "event"
                                      "time"
 [3] "HRV_SD1"
                                      "HRV SD2"
 [5] "HRV_SD1SD2"
                                      "HRV_S"
 [7] "HRV_CSI"
                                      "HRV_CVI"
                                      "HRV_PIP"
 [9] "HRV_CSI_Modified"
[11] "HRV_IALS"
                                      "HRV_PSS"
[13] "HRV_PAS"
                                      "HRV_GI"
[15] "HRV_SI"
                                      "HRV_AI"
[17] "HRV_PI"
                                      "HRV_C1d"
[19] "HRV_C1a"
                                      "HRV_SD1d"
[21] "HRV_SD1a"
                                      "HRV_C2d"
[23] "HRV_C2a"
                                      "HRV_SD2d"
[25] "HRV SD2a"
                                      "HRV Cd"
[27] "HRV_Ca"
                                      "HRV_SDNNd"
[29] "HRV SDNNa"
                                      "HRV ApEn"
[31] "HRV_ShanEn"
                                      "HRV_FuzzyEn"
[33] "HRV_MSEn"
                                      "HRV_CMSEn"
[35] "HRV_RCMSEn"
                                      "HRV_CD"
[37] "HRV_HFD"
                                      "HRV_KFD"
[39] "HRV_LZC"
                                      "HRV_DFA_alpha1"
[41] "HRV_MFDFA_alpha1_Width"
                                      "HRV_MFDFA_alpha1_Peak"
[43] "HRV_MFDFA_alpha1_Mean"
                                      "HRV_MFDFA_alpha1_Max"
[45] "HRV_MFDFA_alpha1_Delta"
                                      "HRV_MFDFA_alpha1_Asymmetry"
[47] "HRV_MFDFA_alpha1_Fluctuation" "HRV_MFDFA_alpha1_Increment"
```

data <- tibble::as_tibble(data)</pre>

```
# * It is very hard to compare the HR as different predictors are on different magnitudes, so we need to normalize them.

time_col <- data$time
```

```
event_col <- data$event
data <- data %>%
    select(-c(time, event)) %>%
    mutate(across(where(is.numeric), scale)) %>%
    mutate(
        time = time_col,
        event = event_col
    )
```

Note now the interpretation of HR is different! For example, if HR=1.16 for the predictor in the univariate model fitted using scaled data, it means that each standard deviation increase is associated with 16% higher risk of event.

```
# For RSF model, we don't need to exclude the missing values
```

Random Survival Forest (RSF)

Variable Selection

The method argument can be set to vh instead for variable hunting, which should be used for problems where the number of variables is substantially larger than the sample size.

```
n_cores <- parallel::detectCores() - 1
cl <- makeCluster(n_cores)
registerDoParallel(cl)

rsf_var_select <- var.select.rfsrc(Surv(time, event) ~ .,
    data = data,
    method = "md",
    seed = 1234,
    ntree = 200,
    parallel = TRUE
) # minimal depth variable selection</pre>
```

```
vars_ranked <- rsf_var_select$topvars</pre>
```

Cross Validation to Select the Best Number of Features

We will use 10-fold cross validation to select the best number of features used in the model.

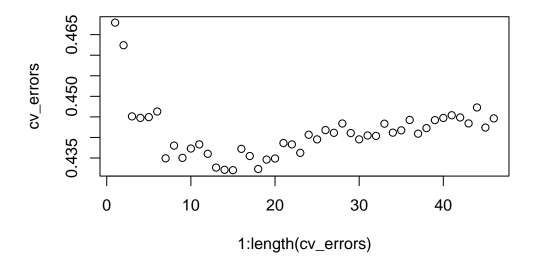
```
set.seed(1234)
folds <- createFolds(data$event, k = n folds) # return indices of folds
cv_errors <- pmclapply(seq(1, length(vars_ranked), by = 1),</pre>

   function(num vars) {
    selected_vars <- vars_ranked[1:num_vars]</pre>
    fold_errors <- sapply(folds, function(fold_idx) {</pre>
        train_data fold <- data[-fold idx, c("time", "event", selected vars)]</pre>
        val_data_fold <- data[fold_idx, c("time", "event", selected_vars)]</pre>
        model <- rfsrc.fast(Surv(time, event) ~ .,</pre>
            data = train_data_fold,
            ntree = 200.
            forest = TRUE
        pred <- predict(model,</pre>
            newdata = val data fold,
            na.action = "na.impute" # * There may be missing values in the

→ dataset

        ) $predicted # define pred has attributes survival(sample_size*time)
 → and predicted(sample_size) for risk
        # Use C-index to measure the performance of the model
        1 - concordance.index(
            pred, # pass risk prediction for first argument
            val_data_fold$time,
            val_data_fold$event
        )$c.index
    })
    mean(fold_errors)
}, title = "Cross Validation to Select the Best Number of Features")
```

```
cv_errors <- as.numeric(cv_errors)
plot(1:length(cv_errors), cv_errors)</pre>
```



```
best_num_vars <- which.min(cv_errors)
vars_selected <- vars_ranked[1:best_num_vars]
print(paste0("The best number of features to retain is ", best_num_vars))</pre>
```

[1] "The best number of features to retain is 15"

Model Fitting

```
data_selected <- data[, c("time", "event", vars_selected)]

# Before formally fitting the model, we can tune the hyperparameters to find:
# 1. optimal mtry (possible split at each node)
# 2. optimal nodesize (minimum size of terminal nodes)
rsf_tuned <- tune.rfsrc(
    Surv(time, event) ~ .,
    data = data_selected,
)</pre>
```

```
rsf_model <- rfsrc(Surv(time, event) ~ .,
    data = data_selected,
    ntree = 500,
    mtry = rsf_tuned$best.mtry,
    nodesize = rsf_tuned$best.nodesize
)</pre>
```

```
# We also fit the full model
rsf_tuned_full <- tune.rfsrc(
    Surv(time, event) ~ .,
    data = data,
)

rsf_model_full <- rfsrc(Surv(time, event) ~ .,
    data = data,
    ntree = 1000,
    mtry = rsf_tuned_full$best.mtry,
    nodesize = rsf_tuned_full$best.nodesize
)</pre>
```

```
# plot.rfsrc?
# plot.variable.rfsrc?
# print.rfsrc?
```