Supplementary Material II: Simulation

Two Stage Design for Prognostic Biomarker Signatures with Survival Endpoint

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This article provides a guidance of the supplementary R functions for the manuscript "Two-Stage Adaptive Design for Prognostic Biomarker Signatures with a Survival Endpoint"

Documentation of Functions

two stage sim()

Description

Conduct simulation for the proposed two-stage design for given scenarios.

Arguments

nn: Total sample size

rate_I: rate parameter in the Exponential Distribution that generates the Time of Event Occurs (default = 0.025)

rate_c: rate parameter in the Exponential Distribution that generates the Time of Censoring (default = 0.025)

beta: the true regression coefficients

mean: parameter in generating the dependent variables (default = 0)

sd: parameter in generating the dependent variables (default = log(1.5))

itrB: number of iterations used for bootstrap or permutation resampling (default = 200)

nfold: number of folds for cross-validation (default = 10)

S: proportion of samples used in first stage (default: 0.5)

alpha_1: level of test in first stage (default: 0.25)

alpha_2: level of test in second stage (default: 0.2)

null_C: The value of C Index under the null hypothesis of interest for bootstrap (default: 0.5)

method: character that takes value of "permutation" or "bootstrap".

rep = 500: total number of iterations for the Monte Carlo Simulation

seedID: seed number for reproducing results (default: NULL)

two stage permutation()

Description

Analyze a given data set using the two-stage design. Inference in the first stage is conducted via permutation.

Arguments

survY: output from surv function in survival package

X: a matrix or data frame of variables of interest, without missing value

itrB: number of iteration used in permutation

seed: seed number for reproducing results (default: NULL)

nfold: number of folds used for cross-validation (default: 10)

S: proportion of samples used in first stage (default: 0.5)

alpha_1: level of test in first stage (default: 0.25)

alpha_2: level of test in second stage (default: 0.2)

printmodel: logical value. TRUE indicates the final signature will be print out (default: FALSE)

two stage bootstrap()

Description

Analyze a given data set using the two-stage design. Inference in the first stage is conducted via bootstrap.

Arguments

survY: output from surv function in survival package

X: a matrix or data frame of variables of interest, without missing value

itrB: number of iteration used in permutation

seed: seed number for reproducing results (default: NULL)

nfold: number of folds used for cross-validation (default: 10)

S: proportion of samples used in first stage (default: 0.5)

alpha_1: level of test in first stage (default: 0.25)

alpha_2: level of test in second stage (default: 0.2)

null_C: The value of C Index under the null hypothesis of interest (default: 0.5)

tuning_C()

Description

Function that helps tune the beta coefficients to hit the target level of C for the alternative hypothesis. For a given vector of beta, only the last coefficient will be tuned and all other coefficients will be fixed.

Arguments

nn: number of samples used as the population (default = 10000)

mean: parameter in generating the dependent variables (default = 0)

sd: parameter in generating the dependent variables (default = log(1.5))

rate_I: rate parameter in the Exponential Distribution that generates the Time of Event Occurs (default = 0.025)

p: number of covariates

initial_beta: a vector of beta coefficients to start with (for p and initial_beta, only of the two arguments is needed)

targetC: target value of C Index of interest

by, start, end: parameters that create a sequence of candidate values for the last coefficient. start: where the sequence starts; end: where the sequence ends; by: the increments for the sequence.

seedID: seed number for reproducing results (default: NULL)

tuning_censoring()

Function that helps tune the distribution of censoring time so that censoring hits the targete proportion.

Arguments

nn: number of samples used as the population (default = 1000)

mean: parameter in generating the dependent variables (default = 0)

sd: parameter in generating the dependent variables (default = log(1.5))

rate_I: rate parameter in the Exponential Distribution that generates the Time of Event Occurs (default = 0.025)

by, start, end: parameters for creating the sequence that can be used to tune the last coefficient. start: where the sequence starts; end: where the sequence ends; by: the increments for the sequence.

p: number of covariates

beta: a vector of beta coefficients

target: target value of censoring proportion

seed: seed number for reproducing results (default: NULL)

.cvFolds()

Lower level function that assigns fold IDs given number of folds.

cvcox()

Lower level function that calculates cross-validated linear predictors from Cox Regression

Examples

Example 2: Simulation Under the Null Hypothesis: C = 0.5

We first illustrate how to use the permutation version

```
source("functions.R")
```

```
##
                                                        se_2 final_validation
            survC threshold early_stop
                                              C_2
## [1,] 0.5825983 0.5324791
                                      0 0.5142684 0.04230755
## [2,] 0.4709064 0.5300727
                                      1
                                                                             0
                                               NA
                                                          NA
                                                                             0
## [3,] 0.4957204 0.5342368
                                      1
                                               NA
                                                          NA
## [4,] 0.4377654 0.5213574
                                      1
                                               NA
                                                          NA
                                                                             0
## [5,] 0.4698146 0.5340279
                                               NA
                                                          NA
                                                                             0
## [6,] 0.5383999 0.5231281
                                      0 0.5715279 0.03922860
                                                                             1
##
             cens
                      cens1
                                cens2
## [1,] 0.5433333 0.5666667 0.5200000
## [2,] 0.4733333 0.4933333 0.4533333
## [3,] 0.4966667 0.5000000 0.4933333
## [4,] 0.4766667 0.4600000 0.4933333
## [5,] 0.4733333 0.4400000 0.5066667
## [6,] 0.4666667 0.4266667 0.5066667
```

As a demonstration, we only specified 5 iterations for the Monte Carlo simulation. Usually it takes >= 500 iterations to evaluate operation characteristics.

To compute type one error rate:

```
mean(result1[,"final_validation"])
```

```
## [1] 0.2
```

To compute probability of early stopping:

```
mean(result1[,"early_stop"])
```

```
## [1] 0.5
```

Example 3: Simulate Samples with the population C Index > 0.5

Generating data with C > 0.5 is less straighthfoward than generating data with C = 0.5. The difficulty comes from hitting the right level of the population C Index and also maintaining the correct proportion of censoring.

We provided two functions \textbf{tuning_C} and \textbf{tuning_censoring} to select the parameters that can generate the targeted level of C.

Step 1: Tune the Regression coefficients with the \textbf{tuning_C} function so that the Population C Index reaches a targeted value.

In this example, we aim at having the C Index to be 0.6. We want to have 5 covariates in the model.

```
## $beta
## [1] 0.2 0.2 0.2 0.2
##
## $beta_j
## [1] 0.83
##
## $C
## [1] 0.6011
##
## $targetC
## [1] 0.6
##
## $mean
## [1] 0
##
## $rate 1
## [1] 0.025
##
## $sd
## [1] 0.4054651
```

The first two outputs yield the values of the beta coefficients that should be used later:

```
tuned_C$beta

## [1] 0.2 0.2 0.2 0.2

tuned_C$beta_j
```

```
## [1] 0.83
```

Step 2: Tune Censoring Distribution with the \textbf{tuned_censor} function to Hit the Targeted Censoring Percentage

Given the beta coefficients produced by \textbf{tuned_C}, We need to tune the rate parameter in the censoring distribution so that 30 percent of the data is censored.

```
## $rate c
## [1] 0.0101
##
## $cpct
## [1] 0.3005
##
## $beta
## [1] 0.20 0.20 0.20 0.20 0.83
##
## $target
## [1] 0.3
##
## $mean
## [1] 0
##
## $rate_1
## [1] 0.025
##
## $sd
## [1] 0.4054651
```

Here we take rate c from the output as the rate parameter for the censoring distribution.

```
tuned_censor$rate_c
```

```
## [1] 0.0101
```

Step 3: Apply the tuned parameters in simulation:

We can use these data to conduct a simulation with the permutation version of the design. In this scenario, the null Hypothesis is C = 0.5. Since the data is generated under C = 0.6, we will be looking at the power of the study.

```
survC threshold early_stop
##
                                             C 2
                                                       se 2 final validation
## [1,] 0.4921614 0.5265356
                                              NA
## [2,] 0.5992083 0.5310401
                                     0 0.5698062 0.03090649
                                                                            1
## [3,] 0.5797136 0.5209427
                                     0 0.5927127 0.03137799
                                                                            1
                                     0 0.5655116 0.03193427
## [4,] 0.5603668 0.5215057
                                                                            1
## [5,] 0.5724786 0.5146318
                                     0 0.5964515 0.02872107
                                                                            1
## [6,] 0.5757391 0.5282800
                                     0 0.5778368 0.03683050
                                                                            1
##
             cens
                      cens1
                                cens2
## [1,] 0.2933333 0.3333333 0.2533333
## [2,] 0.2733333 0.2666667 0.2800000
## [3,] 0.2900000 0.2400000 0.3400000
## [4,] 0.2966667 0.2600000 0.3333333
## [5,] 0.2833333 0.2933333 0.2733333
## [6,] 0.2966667 0.2866667 0.3066667
```

To compute Power:

```
mean(result_P[,"final_validation"])
```

```
## [1] 0.892
```

To compute probability of Early Stopping:

```
mean(result_P[,"early_stop"])
```

```
## [1] 0.082
```

We can use these data to conduct a simulation with the bootstrap version. In this scenario, we can set the null Hypothesis to be C = 0.6. Since the data is generated under C = 0.6, we will be looking at the type one error rate of the study.

```
survC threshold early_stop
                                             C_2
                                                       se_2 final_validation
##
## [1,] 0.6234611 0.6163058
                                     0 0.6304748 0.02927984
                                                                            1
## [2,] 0.6368883 0.6282142
                                     0 0.5979381 0.02954532
                                                                            0
## [3,] 0.5314941 0.5227332
                                                                            0
                                     1
                                              NA
                                                         NA
## [4,] 0.6055463 0.6141665
                                     0 0.5627390 0.03288878
                                                                            0
## [5,] 0.5902689 0.5736166
                                                                            0
                                     1
                                              NA
                                                         NA
## [6,] 0.6422578 0.6437508
                                     0 0.5507143 0.03333810
                                                                            0
##
             cens
                      cens1
                                cens2
## [1,] 0.3366667 0.3866667 0.2866667
## [2,] 0.2766667 0.2600000 0.2933333
## [3,] 0.2800000 0.2733333 0.2866667
## [4,] 0.2800000 0.2666667 0.2933333
## [5,] 0.2966667 0.2733333 0.3200000
## [6,] 0.2900000 0.2933333 0.2866667
```

To compute Type one error rate:

```
mean(result_B[,"final_validation"])
```

```
## [1] 0.018
```

To compute probability of Early Stopping:

```
mean(result_B[,"early_stop"])
```

```
## [1] 0.778
```