# Vignette PlasmodeSim

## 2022-10-19

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Welcome to the vignette about the R package PlasmodeSim. This package is still under development. The goal of this package is to simulate new outcomes for real patients data. This way one can obtain outcomes that follow a model you specify.

## Installing plasmodeSim using remotes

One can easily install the package using remotes, run:

```
install.packages("remotes")
remotes::install_github("GidiusVanDeKamp/PlasmodeSim")
```

## Logistic Regression/ binary outcomes

We start by simulating simple binary variables. In this vignette we use a logisite regression as model, but one could also pick other models that can be implemented as a plpModel. ## Setting up To start we need a plpModel and plpData. For information how to obtain these one can look at: https://ohdsi.github.io/PatientLevelPrediction/articles/BuildingPredictiveModels.html In this documents we load them from a save file:

```
plpResultLogistic <- PatientLevelPrediction::loadPlpResult( "yourpathForPlpResult")
plpData <- PatientLevelPrediction::loadPlpData( "yourPathForPlpData" )</pre>
```

### Example 1 Simulate from a plpModel

In this example we obtain new outcomes following a fitted logistic model. We start from a plpModel, then run predictPlp. At last we generate new outcomes with the function newOutcomes that uses the plpPrediction.

```
plpModelLog <- plpResultLogistic$model

plpPrediction <- PatientLevelPrediction::predictPlp(
    plpModel = plpModelLog,
    plpData = plpData,
    population = plpData$cohorts
)</pre>
```

```
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.201 secs
## Prediction took 0.186 secs
```

When running the function predictPlp it returns some information.

```
newOut <- PlasmodeSim::newOutcomes(
   noPersons = 200,
   props = plpPrediction
)
head(newOut)</pre>
```

```
##
     rowId outcomeCount
## 1
         19
## 2
         26
                         0
                         0
## 3
         29
## 4
         55
                         0
## 5
         67
                         0
                         0
## 6
```

The column called rowId in the output of newOutcomes are the rowId's of patients that are drawn randomly with the same probability, the patients could be drawn multiple times. If a rowId happens to be in the output twice they can have a different outcome, but have the same probability distribution. The function newOutcomes needs a data set that contains the columns rowId and value. The column called value contains the probability of seeing an outcome.

## Example 2 simulation from unfittedmodel

We here we show how to simulate outcomes from an unfitted logistic model. We use the function makeLogisiticModel to specify a logistic model.

```
Parameters <- plpModelLog$model$coefficients
UnfittedParameters <- Parameters
UnfittedParameters[1,1] <- -0.4
UnfittedParameters[3:5,1] <- 0.4
head(UnfittedParameters)
```

```
##
     betas covariateIds
## 1 -0.4
           (Intercept)
## 2
       0.0
                   6003
## 3
       0.4
                    8003
## 4
       0.4
                    9003
       0.4
## 5
                8507001
## 6
      0.0
               28060210
```

For the logistic model it is necessary that the parameters are stored in a dataset with a column called betas and a column called covariateIds. The function makeLogisitcModel makes a plpModel from the specified parameters. The parameters have are given in a data frame with columns called betas and covariateIds. The column called betas has the parameters of the model as numeric values. The columns called covariateIds has its elements stored as a string being '(Intercept)' or a covariateId.

```
plpModelunfitted <- PlasmodeSim::makeLogisticModel(UnfittedParameters)
newprobs <- PatientLevelPrediction::predictPlp(
   plpModel = plpModelunfitted,
   plpData = plpData,
   population = plpData$cohorts
)</pre>
```

```
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.179 secs
## Prediction took 0.184 secs
```

```
newOut <- PlasmodeSim::newOutcomes(
  noPersons = 2000,
  props = newprobs
)
head(newOut)</pre>
```

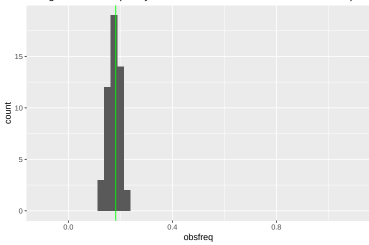
```
##
     rowId outcomeCount
## 1
         3
## 2
         6
                       1
         7
## 3
                       1
## 4
         7
                       1
## 5
         8
## 6
         8
```

## Visual simulations

The function visualOutcome simulates new data and then plots the frequency of the outcome. Right now the function visualOutcome only works for a logistic model. The green line in the plots is the average outcome in the original data set.

```
PlasmodeSim::visualOutcome(
   plpData = plpData,
   noSimulations = 50,
   noPersons = 400,
   parameters = Parameters
)
```

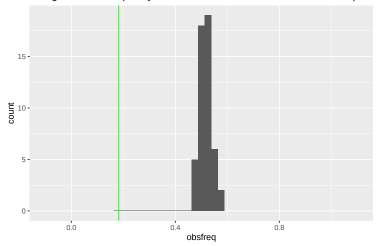
- ## Removing infrequent and redundant covariates and normalizing
  ## Removing infrequent and redundant covariates covariates and normalizing took 0.198 secs
  ## Prediction took 0.172 secs
  - histogram of the frequency of the outcome for 50 simulations with 400 perso



```
PlasmodeSim::visualOutcome(
  plpData = plpData,
  noSimulations = 50,
  noPersons = 400,
  parameters = UnfittedParameters
)
```

## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.192 secs
## Prediction took 0.185 secs





Here we have plotted 50 times the frequency of the outcome for a simulated dataset with 200 people. We can see that the outcome count for the fitted parameters is similar as in the original dataset, but when changing the parameters the outcome count also changes.

## Visual of a specific covariate

Say we are interested in the outcomes of a group with a specific covariate. Here we picked the third covariate in the model to visualise.

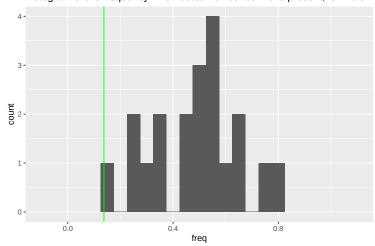
```
covariateIdToStudy<- plpResultLogistic$covariateSummary$covariateId[4]
UnfittedParameters[4,]</pre>
```

```
## betas covariateIds
## 4 0.4 9003
```

```
PlasmodeSim::visualOutcomeCovariateId(
   plpData=plpData,
   studyCovariateId= covariateIdToStudy,
   noSimulations = 20,
   noPersons = 200,
   parameters= UnfittedParameters
)
```

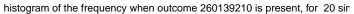
```
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.206 secs
## Prediction took 0.186 secs
```

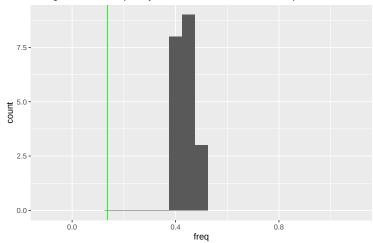
#### histogram of the frequency when outcome 260139210 is present, for 20 sim:



```
PlasmodeSim::visualOutcomeCovariateId2(
   plpData=plpData,
   restrictToCovariateId= covariateIdToStudy,
   noSimulations = 20,
   noPersons= 200,
   parameters= UnfittedParameters
)
```

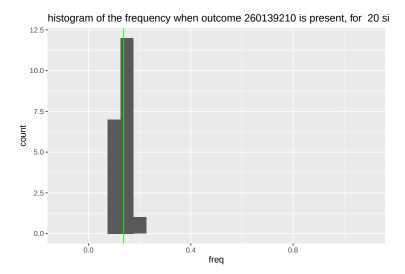
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.189 secs
## Prediction took 0.202 secs





```
PlasmodeSim::visualOutcomeCovariateId2(
   plpData=plpData,
   restrictToCovariateId= covariateIdToStudy,
   noSimulations = 20,
   noPersons= 200,
   parameters= Parameters
)
```

```
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.187 secs
## Prediction took 0.181 secs
```



As one can see visualOutcomeCovariateId and visualOutcomeCovariateId2 are very similiar, they both calculate and plot the frequency for a group with a specific covariate present. The small difference is that visualOutcomeCovariateId filters a newly simulated dataset set to only keep the patients where the covariate is present, and visualOutcomeCovariateId2 only simulates new outcomes for patients that have the covariate present. We see they are almost the identical only visualOutcomeCovariateId2 is spread out less because the groups for calculating the frequency with are larger. Again we see that when picking the fitted parameters the outcome count for patients with a specific covariate is similar as it was in the original data set.

#### Survival times

In this part we will show how to simulate new survival times. For simulating new censored survival times we need more than one probability, we use the baselinehazard, stored in the a plpModel.

#### Loading the plpData

The first step is to load the data where we will simulate new outcomes for. Here we use the package eunomia for accessing some data set.

```
connectionDetails <- Eunomia::getEunomiaConnectionDetails()

Eunomia::createCohorts(
   connectionDetails = connectionDetails,
   cdmDatabaseSchema = 'main',
   cohortDatabaseSchema = 'main',
   cohortTable = 'cohort'
)</pre>
```

```
## Creating cohort: Celecoxib
## |
```

```
## Creating cohort: Diclofenac
## |
                                                                                      1
## Creating cohort: GiBleed
## Creating cohort: NSAIDs
                                                                                      ##
## Cohorts created in table main.cohort
##
     cohortId
                    name
## 1
          1 Celecoxib
           2 Diclofenac
## 2
## 3
            3
                 GiBleed
## 4
            4
                  NSAIDs
##
                                                                                            description
## 1
        A simplified cohort definition for new users of celecoxib, designed specifically for Eunomia.
        A simplified cohort definition for new users ofdiclofenac, designed specifically for Eunomia.
## 3 A simplified cohort definition for gastrointestinal bleeding, designed specifically for Eunomia.
           A simplified cohort definition for new users of NSAIDs, designed specifically for Eunomia.
    count
## 1 1844
## 2
      850
## 3
       479
## 4 2694
databaseDetails <- PatientLevelPrediction::createDatabaseDetails(</pre>
  connectionDetails = connectionDetails,
  cdmDatabaseId = "eunomia",
  cdmDatabaseSchema = 'main',
  cdmDatabaseName = 'Eunomia',
  cohortDatabaseSchema = 'main',
  cohortTable = 'cohort',
  target = 4,
  outcomeDatabaseSchema = 'main',
  outcomeTable = 'cohort',
 outcomeId = 3,
  cdmVersion = 5
)
covariateSettings <- FeatureExtraction::createCovariateSettings(</pre>
  useDemographicsGender = TRUE,
  useDemographicsAgeGroup = TRUE,
  useConditionGroupEraLongTerm = TRUE,
  useDrugGroupEraLongTerm = TRUE,
  endDays = -1,
  longTermStartDays = -365
restrictPlpDataSettings <- PatientLevelPrediction::createRestrictPlpDataSettings(
  studyStartDate = '20000101',
  studyEndDate = '20200101',
 firstExposureOnly = TRUE,
  washoutPeriod = 30
)
```

```
restrictPlpDataSettings <- PatientLevelPrediction::createRestrictPlpDataSettings(
  firstExposureOnly = TRUE,
  washoutPeriod = 30
)
plpData <- PatientLevelPrediction::getPlpData(</pre>
 databaseDetails = databaseDetails,
  covariateSettings = covariateSettings,
 restrictPlpDataSettings = restrictPlpDataSettings
##
     1
                                                                                      1
## Warning: The 'oracleTempSchema' argument is deprecated. Use 'tempEmulationSchema' instead.
## This warning is displayed once every 8 hours.
## Constructing features on server
                                                                                      1
##
## Fetching data from server
## Fetching data took 0.179 secs
```

#### Defining a training set.

Most of the time we split the dataset into training and a test set. In order to prepare the data for fitting the model we have the function MakeTraingSet. What copies features of the function patientLevelPrediction::runPlp. In order to run it we have to create our settings: populationSettings, executeSettings, splitSettings, sampleSettings, featureEngineeringSettings, preprocessSettings. Besides all these settings it also needs the plpData and the outcomeId.

```
populationSettings <- PatientLevelPrediction::createStudyPopulationSettings(</pre>
 binary = TRUE,
 includeAllOutcomes = FALSE,
 firstExposureOnly = FALSE,
  washoutPeriod = 180,
 removeSubjectsWithPriorOutcome = FALSE,
  priorOutcomeLookback = 99999,
 requireTimeAtRisk = TRUE,
 minTimeAtRisk = 1,
 riskWindowStart = 1,
 startAnchor = 'cohort start',
 riskWindowEnd = 7300,
  endAnchor = 'cohort start'
executeSettings <- PatientLevelPrediction::createExecuteSettings(</pre>
  runSplitData = TRUE,
 runSampleData = FALSE,
 runfeatureEngineering = FALSE,
 runPreprocessData = TRUE,
 runModelDevelopment = TRUE,
 runCovariateSummary = TRUE
splitSettings <- PatientLevelPrediction::createDefaultSplitSetting(</pre>
```

```
trainFraction = 0.75,
  splitSeed = 123,
 nfold = 3,
  type = 'stratified'
sampleSettings <- PatientLevelPrediction::createSampleSettings(</pre>
  type = 'none'
)
featureEngineeringSettings <-</pre>
  PatientLevelPrediction::createFeatureEngineeringSettings(
  type = 'none'
)
preprocessSettings <- PatientLevelPrediction::createPreprocessSettings(</pre>
  minFraction = 0,
 normalize = TRUE,
  removeRedundancy = TRUE
TrainingSet <- PlasmodeSim::MakeTraingSet(</pre>
  plpData = plpData,
  executeSettings = executeSettings,
  populationSettings = populationSettings,
  splitSettings = splitSettings,
  sampleSettings = sampleSettings,
  preprocessSettings = preprocessSettings,
  featureEngineeringSettings = featureEngineeringSettings,
  outcomeId = 3
## Outcome is 0 or 1
## seed: 123
## Creating a 25% test and 75% train (into 3 folds) random stratified split by class
## Data split into 656 test cases and 1974 train cases (658, 658, 658)
## Train Set:
## Fold 1 658 patients with 120 outcomes - Fold 2 658 patients with 120 outcomes - Fold 3 658 patients
## 103 covariates in train data
## Test Set:
## 656 patients with 119 outcomes
## Removing 2 redundant covariates
## Normalizing covariates
## Tidying covariates took 0.5 secs
## Train Set:
## Fold 1 658 patients with 120 outcomes - Fold 2 658 patients with 120 outcomes - Fold 3 658 patients
## 101 covariates in train data
## Test Set:
## 656 patients with 119 outcomes
```

#### Fitting the model with censoring

testFraction = 0.25,

We pick the desired model by setting the modelsettings. Then we can run the function fitModelWithCensoring. This function fits two plpModels one for the censoring and one for outcomes, both of the type specified in the modelsettings. It stores these plpModels as a list.

```
modelSettings <- PatientLevelPrediction::setCoxModel()

fitCensor <- PlasmodeSim::fitModelWithCensoring(
    Trainingset = TrainingSet$Train,
    modelSettings = modelSettings
)

## Running Cyclops
## Done.
## GLM fit status: OK
## Creating variable importance data frame
## Prediction took 0.148 secs
## Running Cyclops
## Done.
## GLM fit status: OK
## Creating variable importance data frame
## Prediction took 0.136 secs</pre>
```

#### Generating new outcomes times

Now that we have our model with the censoring specified, we can simulate new outcomes. We call the function simulateSurvivaltimesWithCensoring. It uses the populationSettings for finding the last time that can be included in the outcome times.

```
NewOutcomes <- PlasmodeSim::simulateSurvivaltimesWithCensoring(
   censorModel = fitCensor,
   plpData = plpData,
   population = TrainingSet$Train$labels,
   populationSettings = populationSettings,
   numberToSimulate = 10
)

## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.175 secs
## Prediction took 0.251 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.178 secs
## Prediction took 0.252 secs</pre>
```

```
head(NewOutcomes)
```

```
rowId survivalTime outcomeCount
##
## 1
     425
                   6096
                   7293
                                   0
## 2 1557
## 3 2066
                   2024
                                   0
                                   0
## 4
      664
                   1329
## 5
       48
                   5593
                                   0
## 6
       299
                     18
                                   1
```

Since the censoring model Stores two models as a list one can easily generate uncensored outcomes by using the function simulateSurvivaltimes. One could also use this function for generating censoring times.

```
newdata <- PlasmodeSim::simulateSurvivaltimes(
   plpModel = fitCensor$outcomesModel,
   plpData = plpData,
   numberToSimulate = 10,
   population = TrainingSet$Train$labels,
   populationSettings = populationSettings
)</pre>
```

```
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.181 secs
## Prediction took 0.247 secs
```

## head(newdata)

```
## rowId outcome
## 1 2536 7300
## 2 1882 7300
## 3 1494 7300
## 4 1609 7300
## 5 911 7300
## 6 2610 7300
```

### Defining an unfitted model without censoring

Just as before we can define a model that has not been fitted to the data. We specify a cox model by specifying the two sets of coefficients/parameters and two baseline survival functions.

```
plpModel <- fitCensor$outcomesModel</pre>
coeff <- plpModel$model$coefficients</pre>
survival <- plpModel$model$baselineSurvival$surv</pre>
times <- plpModel$model$baselineSurvival$time</pre>
unfittedmodel <- PlasmodeSim::defineCoxModel(</pre>
  coefficients = coeff,
  baselinehazard = survival,
 timesofbaselinhazard = times,
  featureEngineering = NULL # = NULL is the standard setting.
)
newdata <- PlasmodeSim::simulateSurvivaltimes(</pre>
  plpModel = unfittedmodel,
  plpData = plpData,
  numberToSimulate = 10,
  population = TrainingSet$Train$labels,
  populationSettings = populationSettings
```

## Prediction took 0.176 secs

#### head(newdata)

```
##
     rowId outcome
## 1
                 18
        17
## 2
     1783
               7300
      2600
               7300
## 3
## 4
       964
               7300
## 5
        30
                 81
## 6 1182
               7300
```

## Defining an unfitted model with censoring

There is no function to define an unfitted model with censoring. However this can be done easily by making to cox models and storing them in a list. This elements in this list should have the names <code>censorModel</code> and <code>outcomeModel</code>. In this example we use the unfitted model, specified in the code above, for the outcomes and use the fitted censoring model.

```
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.18 secs
## Prediction took 0.246 secs
## Prediction took 0.182 secs
```

#### head(NewOutcomes)

|   | ## |   | rowld | survivalTime | outcomeCount |
|---|----|---|-------|--------------|--------------|
| : | ## | 1 | 1393  | 71           | 0            |
| : | ## | 2 | 1363  | 36           | 0            |
| : | ## | 3 | 485   | 7300         | 0            |
| : | ## | 4 | 769   | 7300         | 0            |
| : | ## | 5 | 244   | 30           | 0            |
| : | ## | 6 | 614   | 7300         | 0            |

## Adjusting the BaselineSurvival

If one want to get a grip on the outcome count on a specific time one can call the function adjustBaselineSurvival. This can be useful for when one wants multiple data sets that have different parameters, but with the same frequency of outcomes. The function adjustBaselineSurvival changes the base line function of a model such that, for the training data at the specified time the outcome rate is a specified probability. Since this function solves an equation it needs an specified interval to find this solution.

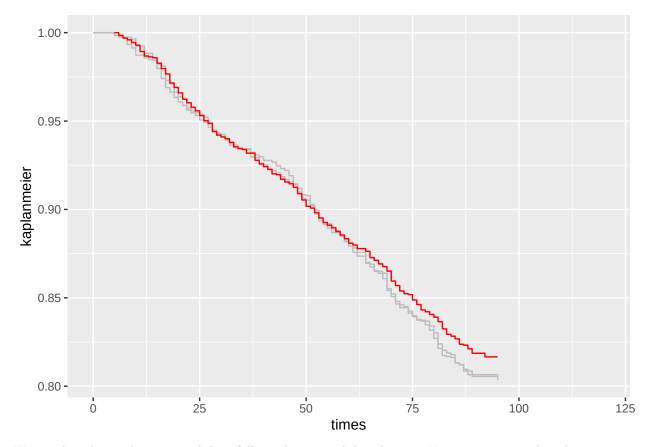
```
adjustedModel <- PlasmodeSim::adjustBaselineSurvival(</pre>
  plpModel = plpModel,
  TrainingSet = TrainingSet$Train,
  plpData = plpData,
  populationSettings = populationSettings,
  timeToFixAt = 3592,
 propToFixWith = 0.87,
  intervalSolution= c(-100,100)
)
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.185 secs
## Prediction took 0.246 secs
NewOutcomes <- PlasmodeSim::simulateSurvivaltimesWithCensoring(</pre>
  censorModel = list(censorModel = fitCensor$outcomesModel,
                     outcomesModel = adjustedModel),
  plpData = plpData,
  population = TrainingSet$Train$labels,
 populationSettings = populationSettings,
  numberToSimulate = 2000
)
## Prediction took 0.186 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.185 secs
## Prediction took 0.246 secs
head(NewOutcomes)
##
    rowId survivalTime outcomeCount
## 1 2367
                    13
## 2 404
                     33
                                   1
## 3 1501
                     14
                                   1
## 4 1545
                     19
                                   1
## 5 1942
                     0
                                   1
## 6 1862
                     18
```

#### Plotting Kaplan Meier estimates

The function kaplanMeierPlot visualizes the Kaplan Meier estimate of a given data set. It works with ggplot. We can easily compare the simulated data sets with the original data by putting them in one plot. For the true data set we set the colour to red.

```
NewOutcomes <- PlasmodeSim::simulateSurvivaltimesWithCensoring(
   censorModel = fitCensor,
   plpData = plpData,
   population = TrainingSet$Train$labels,
   populationSettings = populationSettings,
   numberToSimulate = 1974
)</pre>
```

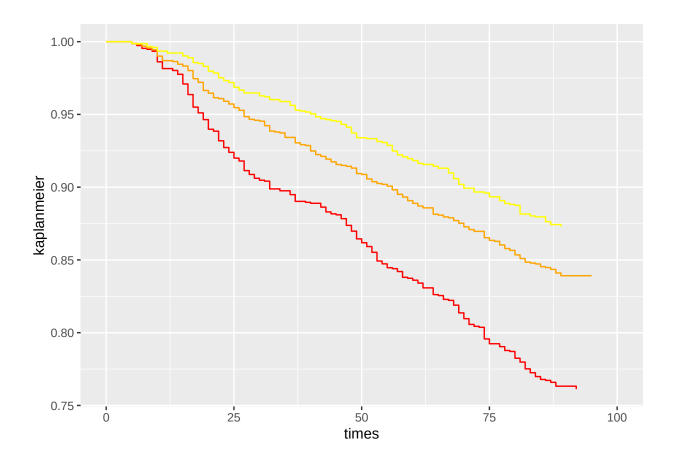
```
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.179 secs
## Prediction took 0.253 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.18 secs
## Prediction took 0.252 secs
NewOutcomes2 <- PlasmodeSim::simulateSurvivaltimesWithCensoring(</pre>
  censorModel = fitCensor,
  plpData = plpData,
  population = TrainingSet$Train$labels,
 populationSettings = populationSettings,
  numberToSimulate = 1974
)
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.178 secs
## Prediction took 0.258 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.179 secs
## Prediction took 0.251 secs
ggplot2::ggplot()+
  PlasmodeSim::KaplanMeierPlot( NewOutcomes )+
  PlasmodeSim::KaplanMeierPlot( NewOutcomes2 )+
  PlasmodeSim::KaplanMeierPlot( TrainingSet$Train$labels, colour = 'red' )+
  ggplot2::xlim(c(0,120))
## Warning: Removed 499 rows containing missing values ('geom_step()').
## Warning: Removed 497 rows containing missing values ('geom_step()').
## Warning: Removed 790 rows containing missing values ('geom_step()').
```



We see that the newly generated data follows the original distribution. However it seems that the outcomes are more frequent in the original dataset.

```
fitCensor$outcomesModel$model$coefficients[2,1] <- 0.5</pre>
fitCensor$outcomesModel$model$coefficients[2,2]
## [1] "8003"
fitCensor$outcomesModel$model$coefficients[3,1] <- 0</pre>
fitCensor$outcomesModel$model$coefficients[3,2]
## [1] "9003"
fitCensor$outcomesModel$model$coefficients[4,1] <- -0.75</pre>
fitCensor$outcomesModel$model$coefficients[4,2]
## [1] "8507001"
numbertosimulate <- 2000
newOut1 <- PlasmodeSim::simulateSurvivaltimesWithCensoringCovariate(fitCensor,</pre>
                                                          plpData,
                                                          TrainingSet$Train$labels,
                                                          populationSettings,
                                                          numbertosimulate,
                                                          8003)
```

```
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.196 secs
## Prediction took 0.224 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.183 secs
## Prediction took 0.218 secs
newOut2 <- PlasmodeSim::simulateSurvivaltimesWithCensoringCovariate(fitCensor,</pre>
                                                        plpData,
                                                        TrainingSet$Train$labels,
                                                        populationSettings,
                                                        numbertosimulate,
                                                        9003)
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.327 secs
## Prediction took 0.196 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.179 secs
## Prediction took 0.195 secs
newOut3 <- PlasmodeSim::simulateSurvivaltimesWithCensoringCovariate(fitCensor,</pre>
                                                        plpData,
                                                        TrainingSet$Train$labels,
                                                        populationSettings,
                                                        numbertosimulate,
                                                        8507001)
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.186 secs
## Prediction took 0.223 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.18 secs
## Prediction took 0.223 secs
ggplot2::ggplot()+
  PlasmodeSim::KaplanMeierPlot( newOut1, colour = 'red')+
  PlasmodeSim::KaplanMeierPlot( newOut2, colour = 'orange')+
  PlasmodeSim::KaplanMeierPlot( newOut3, colour = 'yellow')+
  ggplot2::xlim( c(0,100))
## Warning: Removed 422 rows containing missing values ('geom_step()').
## Warning: Removed 467 rows containing missing values ('geom_step()').
## Warning: Removed 468 rows containing missing values ('geom_step()').
```



#### runPlasmode

## Train Set:

## Test Set:

## 103 covariates in train data

The function runPlasmode returns some new simulated survivaltimes, from a model it fits.

```
runPlas <- PlasmodeSim::runPlasmode(
   plpData = plpData,
   outcomeId = 3,
   populationSettings = populationSettings,
   splitSettings = splitSettings,
   sampleSettings = sampleSettings,
   featureEngineeringSettings = featureEngineeringSettings,
   preprocessSettings = preprocessSettings,
   modelSettings = modelSettings,
   executeSettings = executeSettings,
   numberToSimulate = 5
)

## Outcome is 0 or 1
## seed: 123
## Creating a 25% test and 75% train (into 3 folds) random stratified split by class
## Data split into 656 test cases and 1974 train cases (658, 658)</pre>
```

## Fold 1 658 patients with 120 outcomes - Fold 2 658 patients with 120 outcomes - Fold 3 658 patients

```
## 656 patients with 119 outcomes
## Removing 2 redundant covariates
## Normalizing covariates
## Tidying covariates took 0.625 secs
## Train Set:
## Fold 1 658 patients with 120 outcomes - Fold 2 658 patients with 120 outcomes - Fold 3 658 patients
## 101 covariates in train data
## Test Set:
## 656 patients with 119 outcomes
## Running Cyclops
## Done.
## GLM fit status: OK
## Creating variable importance data frame
## Prediction took 0.134 secs
## Running Cyclops
## Done.
## GLM fit status: OK
## Creating variable importance data frame
## Prediction took 0.145 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.181 secs
## Prediction took 0.249 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.183 secs
## Prediction took 0.248 secs
```

#### runPlas

| ## |   | rowId | survivalTime | ${\tt outcomeCount}$ |
|----|---|-------|--------------|----------------------|
| ## | 1 | 572   | 5484         | 0                    |
| ## | 2 | 1726  | 6611         | 0                    |
| ## | 3 | 419   | 7111         | 0                    |
| ## | 4 | 522   | 27           | 1                    |
| ## | 5 | 425   | 7293         | 0                    |

#### Possible extencions

Here is a list of future extensions to make the package more useful:

- The runPlasmode should have a working analysisId, analysisName and logsettings, like runPlp has.
- one could extend the fit model by adding an option for a Different models for the censoring.
- Take a look at the feature ingeneering in the definecoxmodel function.
- Add more functions that define unfitted models.
- Make it faster by filtering the population on the row ids drawn, before making their outcomes.