# 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 patient 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")
```

## problem

This package is designed to simulated datasets. So that one could test different statistical methods on these data. Beside simulating these new datasets, we implemented some ways to visualise the data.

We want to obtain a data set that follows a certain model, so that one can test different models on this data set. We have an outcome Y and some data  $X_1, X_2, \dots, X_n$  for each patient. We want to generate new Y while leaving the X's as they are. This way we keep some characteristics of the data set, but we do have a true model. What makes it an good set to test a statistical method.

In the first part of this vignette we show how to generate data in the case that Y is binary. We will show how to generate Y's that follow a logistic model. So we have

$$P(Y = 1|X_1 \cdots, X_n) = \frac{1}{1 + \exp(-(\beta_0 + \beta_1 X_1 + \beta_2 X_2 \cdots \beta_3 X_3))}$$

, for some  $\beta$ 's we specify. However the ways of generating new data that follow a different model is also possible, if it is implemented in the plpPredict. So the plpData stores the information about each patient  $Y, X_1, \dots, X_n$ , and the plpModel describes the chance of an outcome from the data from a patient. ## 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 saved file:

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

#### Simulate from a plpModel

In this example we obtain new outcomes following a fitted logistic model. We start from a plpModel, then run predictPlp. after that 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 0.269 secs
## Prediction took 0.188 secs
```

When running the function predictPlp it returns some information.

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

```
## rowId outcomeCount
## 1 1 1 1
```

```
## 2 2 0
## 3 2 0
## 4 3 0
## 5 3 0
## 6 4
```

The column called 'rowId' in the output of newOutcomes contains 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, it can have different outcomes, but follows 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.

#### Simulation from an unfitted model

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
## 5
       0.4
                 8507001
## 6
       0.0
                28060210
```

For the logistic model it is necessary that the parameters are stored in a data set with a column called 'betas' and a column called 'covariateIds'. The function makeLogisiticModel creates a plpModel from the specified parameters. The parameters 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 0.218 secs
## Prediction took 0.201 secs
```

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

##		rowId	outcomeCount
##	1	3	1
##	2	3	0
##	3	4	1
##	4	7	1
##	5	8	1
##	6	8	1

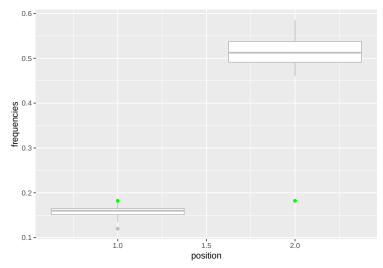
#### Visual simulations

As mentioned before it is important to test on data that is similar to the data that a model will be used for (or trained on). That is why we want to see if the properties from the original set are also present in the test set. We will do this with plots.

We can use the functions frequencyOutcomePlot and frequencyCovariatePlot to plot the frequencies in the new data sets. These functions works together ggplot. We have generated 2000 points. The function calculates multiple frequencies by chopping the data in smaller sets. It also shows the frequency of the outcomes in the plpData with a green dot.

So in the plot below we see on the the outcome frequency of the original dataset with a greendot. On the left we see the frequencies of the data set generated with the fitted model. There are 10 frequencies calculated with groups of 200 patients. On the right we see the same but for the generated data set that comes from an unfitted model. This function makes a random division to calculate the groups. So one might want to use set.seed(). We see that for the fitted model the outcome frequency is similar to the original data set.

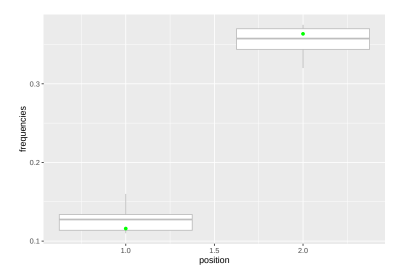
```
ggplot2::ggplot()+
PlasmodeSim::frequencyOutcomePlot(newOutcomesFittedModel, 10, 200, plpData, 1)+
PlasmodeSim::frequencyOutcomePlot(newOutcomesUnfitted, 10, 200, plpData, 2)
```



We are also interested in seeing the fre-

quencies of a specific covariate in the new data set. We do this with the function frequencyCovariatePlot. This function works quite similar, it also chops up the data set. Again we have frequency of the original data set present as a green dot.

```
ggplot2::ggplot()+
PlasmodeSim::frequencyCovariatePlot(newOutcomesFittedModel, 10, 200, 6003, plpData, 1)+
PlasmodeSim::frequencyCovariatePlot(newOutcomesFittedModel, 10, 200, 8003, plpData, 2)
```



## Survival times/ Cox model

In this part we will show how to simulate new survival times. For simulating new censored survival times, we need more than one probability, so we make use of the baseline hazard, 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()</pre>
Eunomia::createCohorts(
  connectionDetails = connectionDetails,
  cdmDatabaseSchema = 'main',
  cohortDatabaseSchema = 'main',
  cohortTable = 'cohort'
## Creating cohort: Celecoxib
##
## Creating cohort: Diclofenac
##
## Creating cohort: GiBleed
##
## Creating cohort: NSAIDs
##
## Cohorts created in table main.cohort
     {\tt cohortId}
##
                     name
## 1
               Celecoxib
            1
## 2
            2 Diclofenac
## 3
            3
                  GiBleed
```

```
## 4
            4
                  NSAIDs
##
                                                                                            description
        A simplified cohort definition for new users of celecoxib, designed specifically for Eunomia.
## 1
        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
      479
## 3
## 4 2694
databaseDetails <- PatientLevelPrediction::createDatabaseDetails(
  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
```

##

```
## Warning: The 'oracleTempSchema' argument is deprecated. Use 'tempEmulationSchema' instead.
## This warning is displayed once every 8 hours.

## Constructing features on server
## |
## Fetching data from server
## Fetching data took 0.212 secs
```

#### Defining a training set.

Most of the time we split the dataset into a training set and a testing set. In order to prepare the data for fitting the model, we have the function MakeTraingSet. This function 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>
 testFraction = 0.25,
 trainFraction = 0.75,
 splitSeed = 123,
 nfold = 3,
 type = 'stratified'
sampleSettings <- PatientLevelPrediction::createSampleSettings(</pre>
 type = 'none'
featureEngineeringSettings <-</pre>
 PatientLevelPrediction::createFeatureEngineeringSettings(
  type = 'none'
```

```
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 0.516 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:
```

preprocessSettings <- PatientLevelPrediction::createPreprocessSettings(</pre>

#### Fitting the model with censoring

## Creating variable importance data frame

## 656 patients with 119 outcomes

minFraction = 0,
normalize = TRUE,

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. They both are of the type specified with 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</pre>
```

```
## Prediction took 0.132 secs
## Running Cyclops
## Done.
## GLM fit status: OK
## Creating variable importance data frame
## Prediction took 0.143 secs
```

#### Generating new outcome 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.

```
NewOutcomesFittedCensorModel <- PlasmodeSim::simulateSurvivaltimesWithCensoring(
   censorModel = fitCensor,
   plpData = plpData,
   population = TrainingSet$Train$labels,
   populationSettings = populationSettings,
   numberToSimulate = 2000
)

## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.183 secs
## Prediction took 0.254 secs</pre>
```

## Removing infrequent and redundant covariates covariates and normalizing took 0.353 secs

## Removing infrequent and redundant covariates and normalizing

#### head(NewOutcomesFittedCensorModel)

## Prediction took 0.251 secs

```
##
    rowId survivalTime outcomeCount
## 1 425
                 7300
## 2 1557
                  48
## 3 2066
                 7293
                                Ω
## 4 664
                 7300
## 5
                 7293
                                0
      48
## 6
     299
                 2683
```

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

```
NewUnfilteredSurvivaltimes <- PlasmodeSim::simulateSurvivaltimes(
   plpModel = fitCensor$outcomesModel,
   plpData = plpData,
   numberToSimulate = 2000,
   population = TrainingSet$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 0.251 secs
```

#### head(NewUnfilteredSurvivaltimes)

```
##
     rowId outcome
## 1
       640
               7300
## 2
       797
               7300
## 3
      2626
               7300
## 4
      770
                 18
## 5
       162
               7300
## 6
       756
                 54
```

#### 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$outcomesModel
coeff <- plpModel$model$coefficients
survival <- plpModel$model$baselineSurvival$surv
times <- plpModel$model$baselineSurvival$time

unfittedmodel <- PlasmodeSim::defineCoxModel(
    coefficients = coeff,
    baselinehazard = survival,
    timesofbaselinhazard = times,
    featureEngineering = NULL # = NULL is the standard setting.
)

NewOutcomesUnfittedModel <- PlasmodeSim::simulateSurvivaltimes(
    plpModel = unfittedmodel,
    plpData = plpData,
    numberToSimulate = 2000,
    population = TrainingSet$Train$labels,
    populationSettings = populationSettings
)</pre>
```

## Prediction took 0.184 secs

#### head(NewOutcomesUnfittedModel)

```
##
     rowId outcome
## 1 1067
              7300
## 2
         6
              7300
## 3 2068
              7300
              7300
## 4
     1832
     1362
## 5
                65
## 6
       252
              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 two Cox models and storing them in a list. The elements in this list should have the names 'censorModel' and

'outcomeModel'. In this example we use the unfitted model, specified in the code above, for the outcomes and use the fitted censoring model.

```
#we can swap outcomes with censoring.
unfittedcensor <- list(censor Model = unfitted model,
                      outcomesModel = fitCensor$outcomesModel)
NewOutcomesUnfittedCensorModel <- PlasmodeSim::simulateSurvivaltimesWithCensoring(
  censorModel = unfittedcensor,
  plpData = plpData,
  population = TrainingSet$Train$labels,
  populationSettings = populationSettings,
  numberToSimulate = 2000
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.209 secs
## Prediction took 0.29 secs
## Prediction took 0.183 secs
head(NewOutcomesUnfittedCensorModel)
##
     rowId survivalTime outcomeCount
## 1
       399
                   7300
## 2 1195
                   7300
                                   0
## 3 1631
                                   0
                   7300
```

#### Adjusting the BaselineSurvival

7300

7300

7300

## 4 1588

492

869

## 5

## 6

If one wants to get a grip on the outcome count on a specific time, one can call the function adjustBaselineSurvival. This can be useful in cases that one wants to obtain 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 in such a way 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 interval to find this solution specified.

0

0

0

```
adjustedModel <- PlasmodeSim::adjustBaselineSurvival(
   plpModel = plpModel,
   TrainingSet = TrainingSet$Train,
   plpData = plpData,
   populationSettings = populationSettings,
   timeToFixAt = 3592,
   propToFixWith = 0.87,
   intervalSolution= c(-100,100)
)</pre>
```

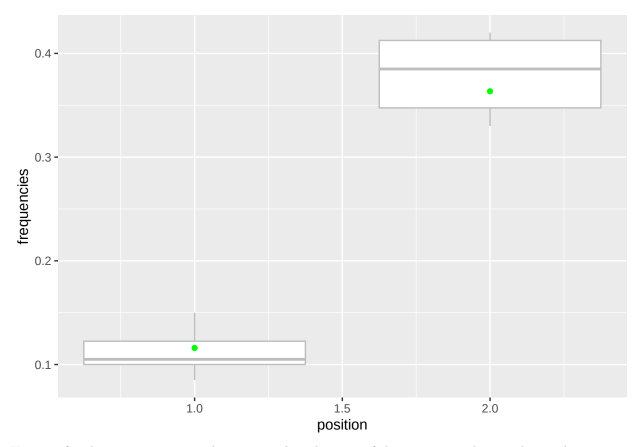
```
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.201 secs
## Prediction took 0.253 secs
```

```
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 0.203 secs
## Prediction took 0.271 secs
head(NewOutcomes)
    rowId survivalTime outcomeCount
##
## 1 1530
                    16
## 2 1041
                                   0
                     25
## 3 1610
                     23
                                   1
## 4 1157
                     23
                                   1
## 5 576
                     26
                                   1
## 6
      913
                     19
                                   1
```

#### Plotting Kaplan Meier estimates

Again we are interested in if the features of the original data set are also in the simulated dataset. We first look at the frequency of the covariates in the simulated data.

```
ggplot2::ggplot()+
PlasmodeSim::frequencyCovariatePlot(NewOutcomesFittedCensorModel, 10, 200, 6003, plpData, 1)+
PlasmodeSim::frequencyCovariatePlot(NewOutcomesFittedCensorModel, 10, 200, 8003, plpData, 2)
```



However for the outcome we are also interested in the time of the outcome. That is why we do not use a boxplot but a Kaplan Meier plot. This type of plot plots the Kaplan Meier estimator that is given by:

$$\hat{S}(t) = \prod_{i:t_i \le t} \left(1 - \frac{d_i}{n_i}\right).$$

Where we have that  $d_i$  is the number of outcomes that happend at time  $t_i$  and  $n_i$  is the number of individuals known to have survived or not yet have been censored upto time  $t_i$ .

The function kaplanMeierPlot visualizes the Kaplan Meier estimate of a given data set. It also 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.

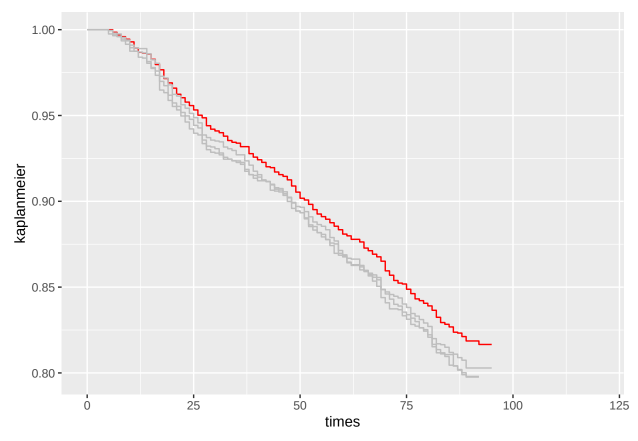
```
toPlotFittedCensorModel <- PlasmodeSim::simulateSurvivaltimesWithCensoring(
  censorModel = fitCensor,
  plpData = plpData,
  population = TrainingSet$Train$labels,
  populationSettings = populationSettings,
  numberToSimulate = 4000
)</pre>
```

## Currently in a tryCatch or withCallingHandlers block, so unable to add global calling handlers. Para ## Currently in a tryCatch or withCallingHandlers block, so unable to add global calling handlers. Para

```
## Removing infrequent and redundant covariates and normalizing
```

## Removing infrequent and redundant covariates covariates and normalizing took 0.199 secs

```
## Currently in a tryCatch or withCallingHandlers block, so unable to add global calling handlers. Para
## Currently in a tryCatch or withCallingHandlers block, so unable to add global calling handlers. Para
## Currently in a tryCatch or withCallingHandlers block, so unable to add global calling handlers. Para
## Prediction took 0.294 secs
## Currently in a tryCatch or withCallingHandlers block, so unable to add global calling handlers. Para
## Currently in a tryCatch or withCallingHandlers block, so unable to add global calling handlers. Para
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.199 secs
## Currently in a tryCatch or withCallingHandlers block, so unable to add global calling handlers. Para
## Currently in a tryCatch or withCallingHandlers block, so unable to add global calling handlers. Para
## Currently in a tryCatch or withCallingHandlers block, so unable to add global calling handlers. Para
## Prediction took 0.289 secs
FittedCensorModel1 <- toPlotFittedCensorModel[1:2000,]
FittedCensorModel2 <- toPlotFittedCensorModel[-(1:2000),]</pre>
FittedCensorModel3 <- NewOutcomesFittedCensorModel
ggplot2::ggplot()+
  PlasmodeSim::KaplanMeierPlot( TrainingSet$Train$labels, colour = 'red' )+
 PlasmodeSim::KaplanMeierPlot(FittedCensorModel1)+
  PlasmodeSim::KaplanMeierPlot(FittedCensorModel2)+
  PlasmodeSim::KaplanMeierPlot(FittedCensorModel3)+
  ggplot2::xlim(c(0,120))
## Warning: Removed 790 rows containing missing values ('geom_step()').
## Warning: Removed 482 rows containing missing values ('geom_step()').
## Warning: Removed 511 rows containing missing values ('geom_step()').
## Warning: Removed 496 rows containing missing values ('geom_step()').
```



Above we see that the newly generated data follows the original distribution. However, it seems that the outcomes are more frequent in the original dataset. We can also generate datasets where all the patients have one specific covariate present. We do this with the function simulateSurvivaltimesWithCensoringCovariate. This function works in a similar way as simulateSurvivaltimesWithCensoring but filters the population to make sure the covariate specified is present.

We are also interested in the influence of a covariate on the survival times. Now we will make the same plots but where the data is filtered such that one specified covariate is present.

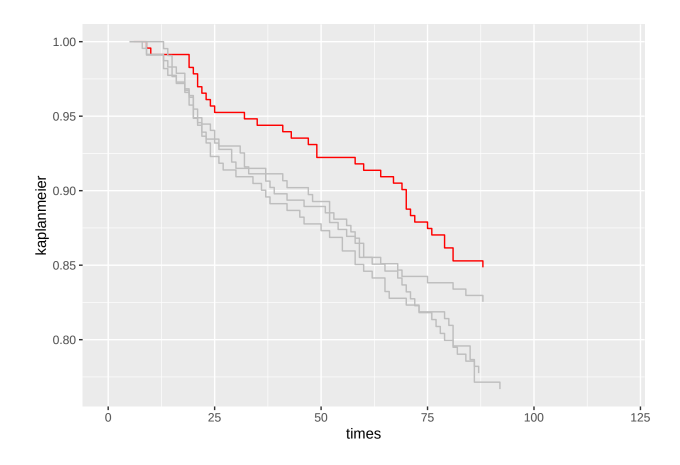
```
ggplot2::ggplot()+
  PlasmodeSim::KaplanMeierPlotFilterCovariate( TrainingSet$Train$labels,6003, plpData, colour = 'red')
  PlasmodeSim::KaplanMeierPlotFilterCovariate( FittedCensorModel1,6003, plpData)+
  PlasmodeSim::KaplanMeierPlotFilterCovariate( FittedCensorModel2,6003, plpData)+
  PlasmodeSim::KaplanMeierPlotFilterCovariate( FittedCensorModel3,6003, plpData)+
  ggplot2::xlim(c(0,120))

## Warning: Removed 87 rows containing missing values ('geom_step()').

## Warning: Removed 70 rows containing missing values ('geom_step()').

## Warning: Removed 95 rows containing missing values ('geom_step()').

## Warning: Removed 85 rows containing missing values ('geom_step()').
```



#### runPlasmode

## 103 covariates in train data

## Test Set:

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

```
runPlas <- PlasmodeSim::runPlasmode(</pre>
  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, 658)
## Train Set:
```

## 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 0.502 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 0.146 secs
## Running Cyclops
## Done.
## GLM fit status: OK
## Creating variable importance data frame
## Prediction took 0.14 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.183 secs
## Prediction took 0.273 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.229 secs
## Prediction took 0.26 secs
```

#### runPlas

##		rowId	survivalTime	outcomeCount
##	1	572	5484	0
##	2	1726	6611	0
##	3	419	7111	0
##	4	522	27	1
##	5	425	7293	0

#### Possible extensions

Following below is a list of suggestions for possible extensions to make the package more useful:

- The runPlasmode should have a working analysisId, analysisName and logsettings, like runPlp has.
- One could extend the fitmodel by adding an option for different models for the censoring.
- Take a look at the feature engineering in the definecoxmodel function.
- Add more functions that define unfitted models.
- Make the functions run faster by filtering the population on the rowids drawn, before making their outcomes.