vignette Survival Times

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first we make sure we have the uptodate package	
remotes::install_github("GidiusVanDeKamp/PlasmodeSim")	

Loading the plpData

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

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

```
## Creating cohort: Celecoxib
## |
## Creating cohort: Diclofenac
## |
## Creating cohort: GiBleed
## |
## Creating cohort: NSAIDs
## |
## Cohorts created in table main.cohort
```

```
## cohortId
## 1 1 Celecoxib
          2 Diclofenac
## 2
## 3
           3 GiBleed
## 4
           4
                 NSAIDs
##
                                                                                           description
        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
      850
## 3
      479
## 4 2694
# Points PatientLevelPredictionPackage to the Eunomia database
# Tells Eunomia to extract the cohort stored with id = 4 as the target cohort
# and cohort with id = 3 as the outcome cohort. The other settings (...Schema)
# tell the database where to look for the target and the outcome cohorts
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
# Use ?FeatureExtraction::createCovariateSettings to see what the options are
# There are a lot...
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
)
```

```
# issue with studyStartDate/studyEndDate
restrictPlpDataSettings <- PatientLevelPrediction::createRestrictPlpDataSettings(
    firstExposureOnly = TRUE,
    washoutPeriod = 30
)

plpData <- PatientLevelPrediction::getPlpData(
    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 0.497 secs</pre>
```

Defining a training set.

For simulating new censored survival times we need more than one probability, we use the baselinehazard, stored in the plpModel, we also need a model for the censoring. First we make the traing set.

```
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,
```

```
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)
## 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 0.658 secs
## Train Set:
```

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

Fitting the model

Test Set:

101 covariates in train data

656 patients with 119 outcomes

splitSeed = 123,
nfold = 3,

To fit the model we make the modelsettings and we run the function fitModelWithCensoring.

```
modelSettings <- PatientLevelPrediction::setCoxModel()
fitCensor <- PlasmodeSim::fitModelWithCensoring(</pre>
```

```
TrainingSet = TrainingSet$Train,
 modelSettings = modelSettings
 # now i have only one model setting
 # should i change this to two seperate settings
)
## Running Cyclops
## Done.
## GLM fit status: OK
## Creating variable importance data frame
## Prediction took 0.188 secs
## Running Cyclops
## Done.
## GLM fit status: OK
## Creating variable importance data frame
## Prediction took 0.164 secs
Generating new outcomes times
population <- PatientLevelPrediction::createStudyPopulation(</pre>
 plpData = plpData,
 outcomeId = 3,
 populationSettings = populationSettings
## Outcome is 0 or 1
NewOutcomes <- PlasmodeSim::simulateSurvivaltimesWithCensoring(</pre>
 censorModel = fitCensor,
 plpData = plpData,
 population = population,
 populationSettings = populationSettings,
 numberToSimulate = 10
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.253 secs
## Prediction took 0.328 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.189 secs
## Prediction took 0.279 secs
head(NewOutcomes)
   rowId survivalTime outcomeCount
##
## 1 2367 18
## 2 500
                  7300
                                  0
## 3 2086
                  20
                                  1
## 4 2279
                 7300
                                  0
## 5 287
                 7300
                                 0
## 6 290
                 5862
```

```
#for simulation the uncensored data

newdata <- PlasmodeSim::simulateSurvivaltimes(
   plpModel = fitCensor$outcomesModel,
   plpData = plpData,
   numberToSimulate = 10,
   population = population,
   populationSettings = populationSettings
)

## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.181 secs
## Prediction took 0.285 secs
head(newdata)</pre>
```

```
## rowId outcome
## 1 1961 7300
## 2 1134 7300
## 3 1720 7300
## 4 2369 7300
## 5 418 7300
## 6 2599 7300
```

Defining an unfitted model

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

Prediction took 0.238 secs

head(newdata) ## rowId outcome ## 1 54 67

```
## 1 54 67

## 2 1229 7300

## 3 552 7300

## 4 2376 7300

## 5 1447 7300

## 6 1709 30
```

Defining an unfitted model with censoring

```
#we can swap outcomes with censoring.
unfittedcensor <- list(censor Model = unfitted model,
                     outcomesModel = fitCensor$outcomesModel)
NewOutcomes <- PlasmodeSim::simulateSurvivaltimesWithCensoring(</pre>
  censorModel = unfittedcensor,
  plpData = plpData,
 population = population,
 populationSettings = populationSettings,
 numberToSimulate = 200
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.219 secs
## Prediction took 0.271 secs
## Prediction took 0.192 secs
head(NewOutcomes)
    rowId survivalTime outcomeCount
## 1 658
                  51
                  7300
## 2 1864
## 3 1317
                 7300
                                  0
## 4 1193
                   86
                                  1
## 5 1518
                  7300
## 6 1883
                  7300
                                  0
```

Adjusting the BaselineSurvival

```
adjustedModel <- PlasmodeSim::AdjustBaselineSurvival(
  plpModel = plpModel,
  TrainingSet = TrainingSet$Train,
  plpData = plpData,
  populationSettings = populationSettings,
  timeTofixat = 3592,</pre>
```

```
proptofixwith = 0.87,
 intervalSolution= c(-100,100)
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.179 secs
## Prediction took 0.254 secs
NewOutcomes <- PlasmodeSim::simulateSurvivaltimesWithCensoring(
 censorModel = list(censorModel = fitCensor$outcomesModel,
                     outcomesModel = adjustedModel),
 plpData = plpData,
 population = population,
 populationSettings = populationSettings,
 numberToSimulate = 2000
)
## Prediction took 0.215 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.192 secs
## Prediction took 0.299 secs
head(NewOutcomes)
##
    rowId survivalTime outcomeCount
## 1 705
                  56
## 2 306
                   14
## 3 1066
                 7300
                                  0
## 4 678
                 7300
                                  0
## 5 532
                  64
                                  0
## 6 1942
                    58
                                  1
```

Plotting Kaplan Meier estimates

the function kaplanMeierPlot visualised the kamplanmeier estimate of a given dataset. It works with ggplot. we can easily compare the simulated data sets with the real dataset 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 = population,
   populationSettings = populationSettings,
   numberToSimulate = 656
)

## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.184 secs
## Prediction took 0.267 secs</pre>
```

Removing infrequent and redundant covariates and normalizing
Removing infrequent and redundant covariates covariates and normalizing took 0.173 secs
Prediction took 0.26 secs

```
NewOutcomes2 <- PlasmodeSim::simulateSurvivaltimesWithCensoring(</pre>
  censorModel = fitCensor,
  plpData = plpData,
  population = population,
  populationSettings = populationSettings,
  numberToSimulate = 656
)
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.176 secs
## Prediction took 0.272 secs
## Removing infrequent and redundant covariates and normalizing
## Removing infrequent and redundant covariates covariates and normalizing took 0.181 secs
## Prediction took 0.265 secs
ggplot2::ggplot()+
 PlasmodeSim::KaplanMeierPlot( NewOutcomes )+
  PlasmodeSim::KaplanMeierPlot( NewOutcomes2 )+
 PlasmodeSim::KaplanMeierPlot( TrainingSet$Test$labels, colour = 'red' )+
 ggplot2::xlim(c(0,120))
## Warning: Removed 227 rows containing missing values ('geom_step()').
## Warning: Removed 231 rows containing missing values ('geom_step()').
## Warning: Removed 283 rows containing missing values ('geom_step()').
   1.00
   0.95 -
 kaplanmeier
   0.90 -
   0.85 -
   0.80 -
                                                                          100
                          25
                                          50
                                                          .
75
           0
                                                                                          125
```

times