# Creating Analysis Specification

## Anthony G. Sena

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# 1 Creating an analysis specification

This walk through will show how to use Strategus to define an analysis specification on an example study using cohorts from the example problem What is the risk of gastrointestinal (GI) bleed in new users of celecoxib compared to new users of diclofenac? as described in the Book Of OHDSI Chapter 12 on Population Level Estimation

## 1.1 Cohorts for the study

To start, we'll need to define cohorts and negative control outcomes to use in our example analysis specification. We've included the cohorts and negative control outcomes in the Strategus package for this example and the code below will load them for use when assembling the analysis specification.

#### 1.1.1 Cohort Definitions & Negative Control Outcomes

This is the list of cohort definitions we will use when assembling the analysis specification for Strategus.

```
kable(cohortDefinitionSet[,c("cohortId", "cohortName")])
```

And the negative control outcomes when performing empirical calibration.

```
kable(ncoCohortSet)
```

## 2 Assembiling HADES modules

The building blocks of the Strategus analysis specification are HADES modules. For purposes of this walk through, a module is a specific analytic task you would like to perform. As shown in the subsequent sections, the high-level pattern for using a module consists of:

- 1. Download the module's settings function.
- 2. Create the module specifications using the settings function(s) from the module
- 3. Compose the analysis pipeline from one or more module settings

## 2.1 CohortGenerator Module Settings

The following code downloads the settings functions from the CohortGeneratorModule which then activates some additional functions we can use for creating the analysis specification. In the analysis specification, we will add the cohort definitions and negative control outcomes to the sharedResources section since these elements may be used by any of the HADES modules. To do this, we will use the createCohortSharedResourceSpecifications and createNegativeControlOutcomeCohortSharedResourceSpecification functions respectively. In addition, we will use the cohortGeneratorModuleSpecifications function to specify the cohort generation settings.

```
source("https://raw.githubusercontent.com/OHDSI/CohortGeneratorModule/v0.0.13/SettingsFunctions.R")

# Create the cohort definition shared resource element for the analysis specification
cohortDefinitionSharedResource <- createCohortSharedResourceSpecifications(
    cohortDefinitionSet = cohortDefinitionSet
)

# Create the negative control outcome shared resource element for the analysis specification
ncoSharedResource <- createNegativeControlOutcomeCohortSharedResourceSpecifications(
    negativeControlOutcomeCohortSet = ncoCohortSet,
    occurrenceType = 'all',
    detectOnDescendants = TRUE)

# Create the module specification
cohortGeneratorModuleSpecifications <- createCohortGeneratorModuleSpecifications(
    incremental = TRUE,
    generateStats = TRUE)</pre>
```

## 2.2 CohortDiagnostics Module Setttings

The following code creates the cohortDiagnosticsModuleSpecifications to run cohort diagnostics on the cohorts in the study.

```
source("https://raw.githubusercontent.com/OHDSI/CohortDiagnosticsModule/v0.0.5/SettingsFunctions.R")
cohortDiagnosticsModuleSpecifications <- createCohortDiagnosticsModuleSpecifications(
    runInclusionStatistics = TRUE,
    runIncludedSourceConcepts = TRUE,
    runOrphanConcepts = TRUE,
    runTimeSeries = FALSE,
    runVisitContext = TRUE,
    runBreakdownIndexEvents = TRUE,
    runIncidenceRate = TRUE,
    runCohortRelationship = TRUE,
    runTemporalCohortCharacterization = TRUE,
    incremental = FALSE)</pre>
```

## 2.3 CohortIncidence Module Setttings

The following code creates the cohortIncidenceModuleSpecifications to perform an incidence rate analysis for the target cohorts and outcome in this study.

```
source("https://raw.githubusercontent.com/OHDSI/CohortIncidenceModule/v0.0.4/SettingsFunctions.R")
library(CohortIncidence)
targets <- list(createCohortRef(id=1, name="Celecoxib"),</pre>
                createCohortRef(id=2, name="Diclofenac"),
                createCohortRef(id=4, name="Celecoxib Age >= 30"),
                createCohortRef(id=5, name="Diclofenac Age >= 30")
);
outcomes <- list(createOutcomeDef(id=1,name="GI bleed", cohortId=3, cleanWindow=9999))
tars <- list(createTimeAtRiskDef(id=1, startWith="start", endWith="end"),</pre>
             createTimeAtRiskDef(id=2, startWith="start", endWith="start", endOffset = 365)
);
analysis1 <- createIncidenceAnalysis(targets = c(1,2,4,5),</pre>
                                      outcomes = c(1),
                                      tars = c(1,2))
irDesign <- createIncidenceDesign(targetDefs = targets,</pre>
                                   outcomeDefs = outcomes,
                                   tars=tars.
                                   analysisList = list(analysis1),
                                   strataSettings = createStrataSettings(
                                     byYear = TRUE,
                                     byGender = TRUE))
cohortIncidenceModuleSpecifications <- createCohortIncidenceModuleSpecifications(</pre>
  irDesign = irDesign$toList()
```

## 2.4 CohortMethod Module Setttings

The following code creates the cohortMethodModuleSpecifications to perform a comparative cohort analysis for this study.

```
library(CohortMethod)
source("https://raw.githubusercontent.com/OHDSI/CohortMethodModule/v0.0.4/SettingsFunctions.R")
negativeControlOutcomes <- lapply(</pre>
 X = ncoCohortSet$cohortId,
 FUN = createOutcome,
 outcomeOfInterest = FALSE,
 trueEffectSize = 1,
  priorOutcomeLookback = 30
outcomesOfInterest <- lapply(</pre>
 X = 3
 FUN = createOutcome,
  outcomeOfInterest = TRUE
outcomes <- append(
  negativeControlOutcomes,
  outcomesOfInterest
tcos1 <- createTargetComparatorOutcomes(</pre>
  targetId = 1,
  comparatorId = 2,
 outcomes = outcomes,
  excludedCovariateConceptIds = c(1118084, 1124300)
)
tcos2 <- createTargetComparatorOutcomes(</pre>
  targetId = 4,
  comparatorId = 5,
 outcomes = outcomes,
  excludedCovariateConceptIds = c(1118084, 1124300)
targetComparatorOutcomesList <- list(tcos1, tcos2)</pre>
covarSettings <- createDefaultCovariateSettings(addDescendantsToExclude = TRUE)</pre>
getDbCmDataArgs <- createGetDbCohortMethodDataArgs(</pre>
  washoutPeriod = 183,
  firstExposureOnly = TRUE,
 removeDuplicateSubjects = "remove all",
 maxCohortSize = 100000,
  covariateSettings = covarSettings
createStudyPopArgs <- createCreateStudyPopulationArgs(</pre>
  minDaysAtRisk = 1,
riskWindowStart = 0,
```

```
startAnchor = "cohort start",
 riskWindowEnd = 30,
  endAnchor = "cohort end"
)
matchOnPsArgs <- CohortMethod::createMatchOnPsArgs()</pre>
fitOutcomeModelArgs <- CohortMethod::createFitOutcomeModelArgs(modelType = "cox")
createPsArgs <- CohortMethod::createCreatePsArgs(</pre>
  stopOnError = FALSE,
  control = Cyclops::createControl(cvRepetitions = 1)
computeSharedCovBalArgs <- CohortMethod::createComputeCovariateBalanceArgs()</pre>
computeCovBalArgs <- CohortMethod::createComputeCovariateBalanceArgs(</pre>
  covariateFilter = FeatureExtraction::getDefaultTable1Specifications())
cmAnalysis1 <- createCmAnalysis(</pre>
  analysisId = 1,
  description = "No matching, simple outcome model",
 getDbCohortMethodDataArgs = getDbCmDataArgs,
 createStudyPopArgs = createStudyPopArgs,
 fitOutcomeModelArgs = fitOutcomeModelArgs
cmAnalysis2 <- createCmAnalysis(</pre>
  analysisId = 2,
  description = "Matching on ps and covariates, simple outcomeModel",
  getDbCohortMethodDataArgs = getDbCmDataArgs,
  createStudyPopArgs = createStudyPopArgs,
  createPsArgs = createPsArgs,
  matchOnPsArgs = matchOnPsArgs,
  computeSharedCovariateBalanceArgs = computeSharedCovBalArgs,
  computeCovariateBalanceArgs = computeCovBalArgs,
  fitOutcomeModelArgs = fitOutcomeModelArgs
cmAnalysisList <- list(cmAnalysis1, cmAnalysis2)</pre>
analysesToExclude <- NULL
cohortMethodModuleSpecifications <- createCohortMethodModuleSpecifications(</pre>
  cmAnalysisList = cmAnalysisList,
  targetComparatorOutcomesList = targetComparatorOutcomesList,
  analysesToExclude = analysesToExclude
```

# 3 Strategus Analysis Specifications

Finally, we will use the various shared resources and module specifications to construct the full set of analysis specifications and save it to the file system in JSON format.

```
analysisSpecifications <- createEmptyAnalysisSpecificiations() %>%
  addSharedResources(cohortDefinitionSharedResource) %>%
  addSharedResources(ncoSharedResource) %>%
  addModuleSpecifications(cohortGeneratorModuleSpecifications) %>%
  addModuleSpecifications(cohortDiagnosticsModuleSpecifications) %>%
  addModuleSpecifications(cohortIncidenceModuleSpecifications) %>%
  addModuleSpecifications(cohortMethodModuleSpecifications)
```

ParallelLogger::saveSettingsToJson(analysisSpecifications, file.path(params\$analysisSettingsPath, param