# Single studies using CohortMethod

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## 1 Introduction

This vignette describes how you can use the CohortMethod package to perform a single new-user cohort study. We will walk through all the steps needed to perform an examplar study, and we have selected the well-studied topic of the effect of coxibs versus non-selective NSAIDs on GI bleeding-related hospitalization.

## 2 Installation instructions

Before installing the CohortMethod package make sure you have Java and RTools installed. Java can be downloaded from www.java.com. RTools can be downloaded from CRAN.

The CohortMethod package is maintained in a Github repository, and has dependencies on other packages in Github. All these packages can be downloaded and installed from within R using the devtools package:

```
install.packages("devtools")
library(devtools)
install_github("ohdsi/SqlRender")
install_github("ohdsi/DatabaseConnector")
install_github("ohdsi/Cyclops")
install_github("ohdsi/CohortMethod")
```

Once installed, you can use library(CohortMethod) to load the package.

## 3 Data extraction

The first step in running the CohortMethod is extracting all necessary data from the server containing the data in Common Data Model format.

## 3.1 Configuring the connection to the server

We need to tell R how to connect to the server where the data is. CohortMethod uses the DatabaseConnector package, which provides the createConnectionDetails function. Type ?createConnectionDetails for the specific settings required for the various database management systems (DBMS). For example, one might connect to a PostgreSQL database using this code:

The last two lines define the cdmSchema and resultSchema variables, which we'll use later to tell R where the data in CDM format lives, and where we want to write intermediate and result tables.

## 3.2 Preparing the exposures and outcome(s)

We need to define the exposures and outcomes for our study. We do this by writing SQL statements against the OMOP Common Data Model that populates a table of events we are interested in. For our example study, we've created a file called *coxibVsNonselVsGiBleed.sql* with the following contents:

```
/***********
File coxibsVsNonselVsGiBleed.sql
*************
USE @cdmSchema;
IF OBJECT_ID('@resultsSchema.dbo.coxibVsNonselVsGiBleed', 'U') IS NOT NULL
 DROP TABLE @resultsSchema.dbo.coxibVsNonselVsGiBleed;
CREATE TABLE @resultsSchema.dbo.coxibVsNonselVsGiBleed (
 cohort_definition_id INT,
   cohort_start_date DATE,
   cohort_end_date DATE,
   subject id BIGINT
   );
INSERT INTO @resultsSchema.dbo.coxibVsNonselVsGiBleed (
   cohort_definition_id,
   cohort start date,
   cohort_end_date,
   subject id
```

```
SELECT 1, -- Exposure
   drug_era_start_date,
   drug_era_end_date,
   person id
FROM drug era
WHERE drug_concept_id = 1118084; -- celecoxib
INSERT INTO @resultsSchema.dbo.coxibVsNonselVsGiBleed (
    cohort definition id,
    cohort_start_date,
    cohort_end_date,
    subject_id
SELECT 2, -- Comparator
   drug_era_start_date,
    drug_era_end_date,
   person_id
FROM drug era
WHERE drug_concept_id = 1124300; --Diclofenac
INSERT INTO @resultsSchema.dbo.coxibVsNonselVsGiBleed (
    cohort definition id,
    cohort_start_date,
    cohort_end_date,
    subject_id
SELECT 3, -- Outcome
    condition_start_date,
    condition_end_date,
    condition_occurrence.person_id
FROM condition_occurrence
INNER JOIN visit_occurrence
    ON condition_occurrence.visit_occurrence_id = visit_occurrence.visit_occurrence_id
WHERE condition_concept_id IN (
        SELECT descendant_concept_id
        FROM concept_ancestor
        WHERE ancestor_concept_id = 192671 -- GI - Gastrointestinal haemorrhage
    AND visit_occurrence.place_of_service_concept_id IN (9201, 9203);
```

This is parameterized SQL which can be used by the SqlRender package. We use parameterized SQL so we do not have to pre-specify the names of the CDM and result schemas. That way, if we want to run the SQL on a different schema, we only need to change the parameter values, we don't have to change the SQL code. By also making use of SqlRender's translation functionality, we can make sure the SQL code can be run in many different environments.

```
sql <- readSql("coxibVsNonselVsGiBleed.sql")
sql <- renderSql(sql, cdmSchema = cdmSchema, resultsSchema = resultsSchema)$sql
sql <- translateSql(sql, targetDialect = connectionDetails$dbms)$sql
connection <- connect(connectionDetails)
executeSql(connection, sql)</pre>
```

In this code, we first read the SQL from the file into memory. In the next line, we replace the two parameter

names with the actual values. We then translate the SQL into the dialect appropriate for the DBMS we already specified in the connectionDetails. Next, we connect to the server, and submit the rendered and translated SQL.

If all went well, we now have a table with the events of interest. We can see how many events per type:

### 3.3 Extracting the data from the server

Now we can tell CohortMethod to define the cohorts based on our events, and extract all necessary data for our analysis:

```
# Get all NSAID Concept IDs for exclusion:
sql <- paste("SELECT concept_id",</pre>
             "FROM concept_ancestor",
             "INNER JOIN concept",
             "ON descendant_concept_id = concept_id",
             "WHERE ancestor_concept_id = 21603933")
nsaids <- querySql(connection, sql)</pre>
nsaids <- nsaids$CONCEPT_ID</pre>
 #Load data:
cohortData <- getDbCohortData(connectionDetails,</pre>
                               cdmSchema = cdmSchema,
                               resultsSchema = resultsSchema,
                               targetDrugConceptId = 1,
                               comparatorDrugConceptId = 2,
                               indicationConceptIds = "",
                               washoutWindow = 183,
                               indicationLookbackWindow = 183,
                               studyStartDate = "",
                               studyEndDate = "",
                               exclusionConceptIds = nsaids,
                               outcomeConceptIds = 3,
                               outcomeConditionTypeConceptIds = "",
                               exposureSchema = resultsSchema,
                               exposureTable = "coxibVsNonselVsGiBleed",
                               outcomeSchema = resultsSchema,
                               outcomeTable = "coxibVsNonselVsGiBleed",
                               useCovariateDemographics = TRUE,
                               useCovariateConditionOccurrence = TRUE,
```

```
useCovariateConditionOccurrence365d = TRUE,
                             useCovariateConditionOccurrence30d = TRUE,
                             useCovariateConditionOccurrenceInpt180d = TRUE,
                             useCovariateConditionEra = TRUE,
                             useCovariateConditionEraEver = TRUE,
                             useCovariateConditionEraOverlap = TRUE,
                             useCovariateConditionGroup = TRUE,
                             useCovariateDrugExposure = TRUE,
                             useCovariateDrugExposure365d = TRUE,
                             useCovariateDrugExposure30d = TRUE,
                             useCovariateDrugEra = TRUE,
                             useCovariateDrugEra365d = TRUE,
                             useCovariateDrugEra30d = TRUE,
                             useCovariateDrugEraEver = TRUE,
                             useCovariateDrugEraOverlap = TRUE,
                             useCovariateDrugGroup = TRUE,
                             useCovariateProcedureOccurrence = TRUE,
                             useCovariateProcedureOccurrence365d = TRUE,
                             useCovariateProcedureOccurrence30d = TRUE,
                             useCovariateProcedureGroup = TRUE,
                             useCovariateObservation = TRUE,
                             useCovariateObservation365d = TRUE.
                             useCovariateObservation30d = TRUE,
                             useCovariateObservationBelow = TRUE,
                             useCovariateObservationAbove = TRUE,
                             useCovariateObservationCount365d = TRUE,
                             useCovariateConceptCounts = TRUE,
                             useCovariateRiskScores = TRUE,
                             useCovariateInteractionYear = FALSE,
                             useCovariateInteractionMonth = FALSE,
                             excludedCovariateConceptIds = nsaids,
                             deleteCovariatesSmallCount = 100)
cohortData
```

```
#> CohortData object
#>
#> Treatment concept ID: 1
#> Comparator concept ID: 2
#> Outcome concept ID(s): 3
```

There are a lot of parameters, but they are all documented in the CohortMethod manual. In short, we're pointing the function to the table created earlier and indicate which concept IDs in that table identify the target, comparator and outcome. We further instruct that people with prior exposure to any NSAID should be excluded, and that many different covariates should be constructed, including covariates for all conditions, drug exposures, and procedures that were found on or before the index date.

All data about the cohorts, outcomes, and covariates are extracted from the server and stored in the cohortData object. This object uses the package ff to store information in a way that ensures R does not run out of memory, even when there is lots of data.

We can use the generic summary() function to view some more information of the data we extracted:

```
summary(cohortData)
```

```
#> CohortData object summary
#>
#> Treatment concept ID: 1
#> Comparator concept ID: 2
#> Outcome concept ID(s): 3
#>
#> Treated persons: 11878
#> Comparator persons: 48415
#>
#> Outcome counts:
     Event count Person count
#> 3
            2254
                          1516
#>
#> Covariates:
#> Number of covariates: 22023
#> Number of non-zero covariate values: 39204318
```

#### 3.3.1 Saving the data to file

Creating the cohortData file can be expensive, and it is probably a good idea to save it for future sessions. Because cohortData uses ff, we cannot use R's regular save function. Instead, we'll have to use the saveCohortData() function:

```
saveCohortData(cohortData, "coxibVsNonselVsGiBleed")
```

We can use the loadCohortData() function to load the data in a future session.

## 4 Propensity scores

The CohortMethod can use propensity scores to adjust for potential confounders. Instead of the traditional approach of using a handfull of predefined covariates, CohortMethod typically uses thousands to millions of covariates that are automatically constructed based on conditions, procedures and drugs in the records of the subjects.

## 4.1 Fitting a propensity model

We can fit a propensity model using the covariates constructed by the getDbCohortData() function:

```
ps <- createPs(cohortData, outcomeConceptId = 3)</pre>
```

The createPs() function uses the Cyclops package to fit a large scale regularized logistic regression. Note that We have to tell createPs what the outcomeConceptId is for which we will use the model so it can remove subjects who had the outcome prior to index date before fitting the model.

To fit the propensity model, Cyclops needs to know the hyperparameter value. By default Cyclops will use cross-validation to estimate the optimal hyperparameter. However, be aware that this can take a really long time. You can use the prior and control parameters of the createPs() to specify Cyclops' behaviour, including using multiple threads to speed up the cross-validation.

## 4.2 Propensity score diagnostics

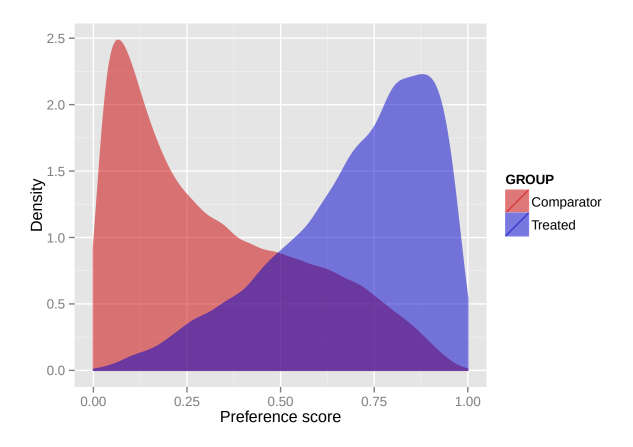
We can compute the Area Under the Receiver-Operator curve:

```
computePsAuc(ps)
```

```
#> [1] 0.8693535
```

We can also plot the propensity score distribution, although we prefer the preference score distribution:

```
plotPs(ps, scale = "preference")
```



It is also possible to inspect the propensity model itself:

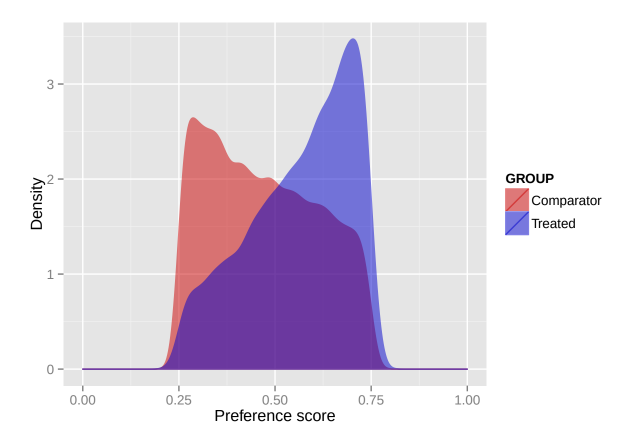
```
propensityModel <- getPsModel(ps, cohortData)
head(propensityModel)</pre>
```

```
#>
        coefficient
                              id
                                                                covariateName
#> 1630
         -3.4137454
                     1150871503
                                ... with cohort index:
                                                         1150871-Misoprostol
#> 1432
        -1.2473805
                              12
                                                            Age group: 10-14
#> 2055
          1.1742889 40664232701 ...IC TREATMENT VISIT (AS PART OF CONTRACT)
         -1.1426356
#> 1
                              13
                                                            Age group: 15-19
#> 1718 -1.1110532 19102773402 ...in 5 MG/ML Ophthalmic Solution [Vigamox]
#> 347
          0.8423181
                           2007
                                                            Index year: 2007
```

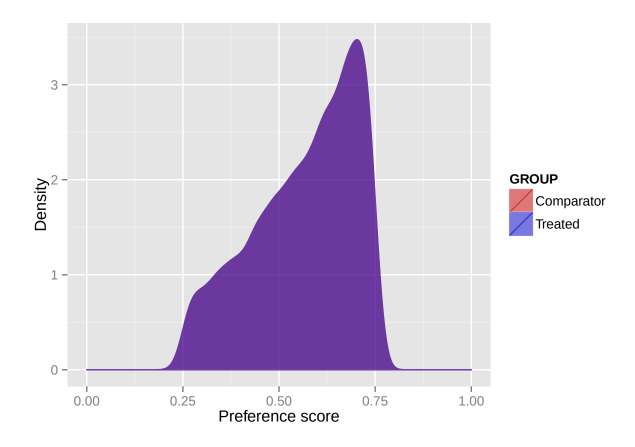
## 4.3 Using the propensity score

We can use the propensity scores to trim, match, or stratify our population. For example, one could first trim to equipoise, meaning only subjects with a preference score between 0.25 and 0.75 are kept:

```
psTrimmed <- trimByPsToEquipoise(ps)
plotPs(psTrimmed, ps, scale = "preference")</pre>
```



We can then match subjects based on propensity scores:

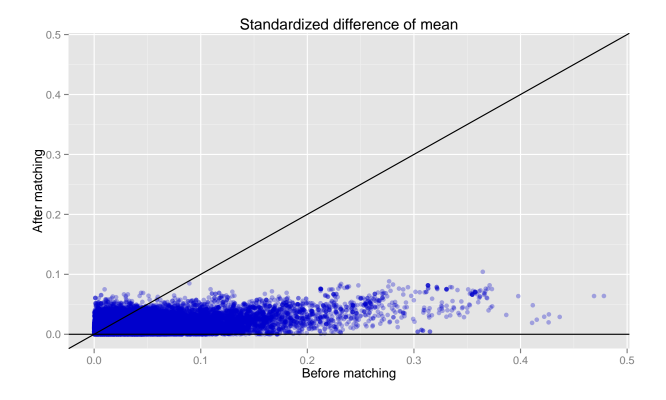


## 4.4 Evaluating covariate balance

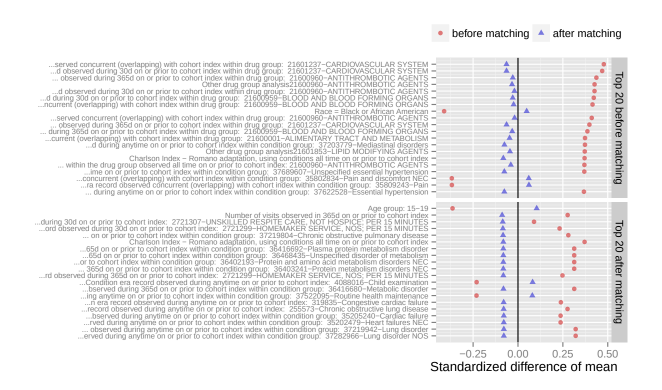
To evaluate whether our use of the propensity score is indeed making the two cohorts more comparable, we can compute the covariate balance before and after trimming, matching, and/or stratifying:

balance <- computeCovariateBalance(strata, cohortData, outcomeConceptId = 3)</pre>

plotCovariateBalanceScatterPlot(balance)



plotCovariateBalanceOfTopVariables(balance)



## 5 Fitting the outcome model

The next step is fitting the outcome model. In theory we could fit an outcome model without using the propensity scores:

In this example we're fitting an outcome model using a Cox regression. The risk window is defined as time of exposure +30 days.

But of course we want to make use of the trimming and matching done on the propensity score:

Note that we define the subpopulation to be only those in the *strata* object, which we created earlier by trimming to equipoise and matching on propensity score. We also now use a stratified Cox model, conditioning on the propensity score match sets.

One final refinement would be to use the same covariates we used to fit the propensity model to also fit the outcome model. This way we are more robust against misspecification of the model, and more likely to remove bias. For this we use the regularized Cox regression in the Cyclops package. (Note that the treatment variable is automatically excluded from regularization.)

We can create the Kaplan-Meier plot:

## plotKaplanMeier(outcomeModel)

We can inspect more details of the outcome model:

```
summary(outcomeModel)
coef(outcomeModel)
confint(outcomeModel)
```

We can also see the covariates that ended up in the outcome model:

```
fullOutcomeModel <- getOutcomeModel(outcomeModel, cohortData)
head(fullOutcomeModel)</pre>
```