Using SelfControlledCohort

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Contents

1	Tables used in the analysis	3
2	A first example	3
3	Custom exposure-outcome pairs	5
4	Running multiple analyses	6
5	Correcting bias with EmpiricalCalibration	8

The Self Controlled Cohort method measures the association between an exposure and an outcome by comparing the number of outcomes during an unexposed time at risk to the number of outcomes during an exposed time at risk. The method is called "Self-Controlled" because each individual in the study contributes person-time to both the exposed and unexposed cohorts. Since each person contributes both exposed and unexposed time to the study only persons who experience the exposure can be used. The inputs to the analysis are the exposures and outcomes of interest, the unexposed time at risk and exposed time at risk dates for each person, and various analysis parameter settings that will be discussed in more detail.

There are seven time points to consider when defining this analysis.

For each exposure-outcome pair the following statistics are calculated:

- Number of persons with the exposure
- Total number of exposures (A person can experience the exposure multiple times.)
- Number of outcomes during the exposed time at risk
- Number of outcomes during the unexposed time at risk
- Total exposed person-time at risk
- Total unexposed person-time at risk
- Incidence rate ratio (incidence rate during exposed time)/(incidence rate during unexposed time) -
- 95% confidence interval for the incidence rate ratio
- The log of the incidence rate ratio The standard error of the log incidence rate ratio

Each person's exposure start date is taken as the index date or $Time\ 0$. The unexposed risk window is defined prior to Time 0 and represents a duration of time when the person was not exposed but could have experienced the outcome. The exposed risk window is defined as a period of time after the Time 0 and represents a duration of time when the person was exposed and could have experienced the outcome. Both

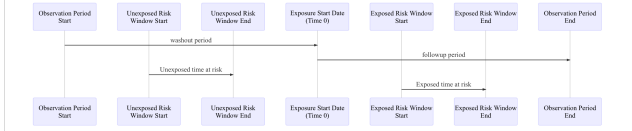


Figure 1: Figure: example exposure windows

of these time windows must be contained within a continuous period of observation. The details about when the risk windows start and end can be adjusted to meet the analysis specifications.

The time between the start of the observation period and Time 0 is called the washout period and a minimum allowed washout period can be set when defining the analysis. Similarly the time between Time 0 and the observation period end date is the followup period. A minimum followup period can also be set.

A common type of exposure is exposure to a drug but an exposure could be any set of characteristics that are captured in your data. We will look at a few different ways to define exposures.

1 Tables used in the analysis

There are two primary tables needed for the self controlled cohort analysis along with a few supporting tables that are part of the OMOP Common Data Model.

Exposure Table

Field	Data type
person_id	Integer
e xposure_concept_id	Integer
e xposure_start_date	Date
$exposure_end_date$	Date

Outcome Table

Field	Data type
person_id	Integer
outcome_concept_id	Integer
outcome_start_date	Date

These tables can be created manually or standard CDM tables can be used. For example by default the drug_era is used as the exposure_table and the condition_era table is used as the outcome_table. Every exposure-outcome combination in these tables will be included in the analysis unless otherwise specified. The additional CDM tables that are required for this analysis are observation_period, necessary for observation start and end dates, and person, necessary for year of birth.

2 A first example

The SelfControlledCohort package makes it easy to calculate the association between every drug and every condition in a CDM database. The Eunomia package provides a mini CDM that runs entirely within R that we can use for examples.

```
library(dplyr)
result$estimates %>%
  arrange(desc(irr)) %>%
  head()
         exposureId outcomeId numPersons numExposures numOutcomesExposed numOutcomesUnexposed timeAtRis
#> 1.200
             757627
                        378419
                                         4
                                                       4
                                                                           4
                                                                                                 0
#> 1.281
            1102527
                       4237458
                                         2
                                                       2
                                                                                                 0
                                                                                                          0.0
```

#> 1.467	1124957	4237458	1	1	1	0
<i>#> 1.671</i>	1177480	4134304	3	3	1	0
<i>#> 1.691</i>	1307863	313217	<i>5</i>	5	5	0
<i>#> 1.753</i>	1310149	313217	5	5	5	0

0.0

0.0

0.0

The result of our analysis is a dataframe with one row per exposure-outcome pair along with all the relevant statistics. Let's interpret the results for amoxicillin exposure (concept ID 1713332) and the outcome of Chronic sinusitis (concept ID 257012)

```
example <- result$estimates %>%
  filter(exposureId == 1713332, outcomeId == 257012)

example %>%
  tidyr::gather() %>%
  mutate(value = format(round(value,1), scientific = F)) %>%
  rename(column = key) %>%
  knitr::kable(align = c("lr"))
```

column	value
exposureId	1713332.0
outcomeId	257012.0
numPersons	4.0
numExposures	4.0
numOutcomesExposed	0.0
${\bf numOutcomes Unexposed}$	1.0
timeAtRiskExposed	0.0
timeAtRiskUnexposed	185.0
irr	0.0
irrLb95	0.0
irrUb95	658680.7
logRr	-Inf
seLogRr	Inf

We can see that 4 were exposed to amoxicillin. When we add up all the time before the exposure we get 184.9609856 person-years. Similarly when we add up all the time after the exposure, the "Exposed time at risk", we get 0.0109514 person-years.

The incidence of Chronic sinusitis during the "Unexposed time at risk" is

$$\frac{224\ events}{45796.0\ person\ years}=0.00489$$

The incidence of Chronic sinusitis during the "Exposed time at risk" is

$$\frac{65\ events}{277.4\ person\ years} = 0.234$$

The incidence rate ratio is

$$\frac{234}{0.00489} = 47.9$$

with a 95% confidence interval of [35.8, 63.4] See vignette ("rateratio.test", package = "rateratio.test") for method details.

3 Custom exposure-outcome pairs

In addition to using all drugs as exposures and all conditions as outcomes we can come up with much more specific definitions of exposures and outcomes using cohorts. A cohort is a set of persons who satisfy one or more inclusion criteria for a duration of time. We can define cohorts using Atlas or by writing SQL code. If we use Atlas then the cohort will be stored in the Atlas "results" schema associated with your CDM database. If you want to write SQL then you will need write access to a schema in your CDM database.

We will simulate using cohorts created in Atlas by using the pre-built cohorts in Eunomia.

Suppose we are interested in the exposure of NSAID (cohort #4) and the outcome of GiBleed (cohort #3). Even though this is a drug-condition pair these cohorts could be defined using combinations of data elements from any domain.

```
result$estimates %>%
  filter(exposureId == 4, outcomeId == 3) %>%
  tidyr::gather() %>%
  mutate(value = format(round(value,1), scientific = F)) %>%
  rename(column = key) %>%
  knitr::kable(align = c("lr"))
```

column value

In this case the risk of being in the GI Bleed cohort before entering the exposure cohort is zero which means our rate ratio is infinite.

Even though this example uses a drug-condition pair for the exposure and the outcome, it demonstrates how we can use any cohorts created in Atlas as exposures and outcomes in a self controlled cohort study.

Let's demonstrate custom exposure outcome pairs using SQL by asking "Do patients tend to get more measurements in the year after a condition diagnosis than the year before?"

```
con <- DatabaseConnector::connect(connectionDetails)
#> Connecting using SQLite driver
```

```
-- create outcome table

create table measurement_cohort as

select

person_id as subject_id,

1 as cohort_definition_id, -- treat any measurement as the same outcome

measurement_date as cohort_start_date,

measurement_date as cohort_end_date

from measurement
```

```
-- create exposure table

create table condition_cohort as

select

person_id as subject_id,

2 as cohort_definition_id, -- treat any condition as the same exposure

condition_start_date as cohort_start_date,

condition_end_date as cohort_end_date

from condition_occurrence
```

Using the riskWindow arguments we can set the time at risk to one year before and one year after each condition exposure.

```
result$estimates %>%
  tidyr::gather() %>%
  mutate(value = format(round(value,1), scientific = F)) %>%
  rename(column = key) %>%
  knitr::kable(align = c("lr"))
```

Indeed it does appear that in Eunomia patients tend to get more tests after a diagnosis than before.

4 Running multiple analyses

The SelfControlledCohort package supports performing multiple analyses at once and provides functions to specify the details of each analysis to be performed.

Start by creating an sccAnalysis object for each analysis type to be performed. We will create one analysis that uses 30 day exposure windows and another with 365 day exposure windows.

```
sccArgs1 <- createRunSelfControlledCohortArgs(firstExposureOnly = TRUE,</pre>
                                                firstOutcomeOnly = TRUE,
                                                minAge = "",
                                                maxAge = "",
                                                studyStartDate = "",
                                                studyEndDate = "",
                                                addLengthOfExposureExposed = TRUE,
                                                riskWindowStartExposed = 1,
                                                riskWindowEndExposed = 30,
                                                addLengthOfExposureUnexposed = TRUE,
                                                riskWindowEndUnexposed = -1,
                                                riskWindowStartUnexposed = -30,
                                                hasFullTimeAtRisk = FALSE,
                                                computeTarDistribution = TRUE,
                                                washoutPeriod = 0,
                                                followupPeriod = 0)
sccArgs2 <- createRunSelfControlledCohortArgs(firstExposureOnly = TRUE,</pre>
                                                firstOutcomeOnly = TRUE,
                                                minAge = "",
                                                maxAge = "",
                                                studyStartDate = "",
                                                studyEndDate = "",
                                                addLengthOfExposureExposed = TRUE,
                                                riskWindowStartExposed = 1,
                                                riskWindowEndExposed = 365,
                                                addLengthOfExposureUnexposed = TRUE,
                                                riskWindowEndUnexposed = -1,
                                                riskWindowStartUnexposed = -365,
```

Then create a list with all exposure outcome pairs to be analyzed. These can be concept IDs if you are using the condition_occurrence and drug_era tables or cohort IDs if you are using a cohort table for exposures and outcomes.

The total number of rate ratios will be length(sccAnalysisList) * length(exposureOutcomesList) since every analysis will be executed for every exposure-outcome pair.

```
results <- runSccAnalyses(connectionDetails,
                         cdmDatabaseSchema = "main",
                         exposureTable = "cohort",
                         outcomeTable = "cohort",
                         outputFolder = "./SelfControlledCohortOutput",
                         sccAnalysisList = sccAnalysisList,
                         exposureOutcomeList = exposureOutcomeList,
                         analysisThreads = 1,
                         computeThreads = 1)
#> *** Running multiple analysis ***
summarizeAnalyses(results, "./SelfControlledCohortOutput")
#> [1] exposureId outcomeId
                                              \it numPersons
                                                                   numExposures
                                              p10TxTime
#> [10] sdTxTime
                           minTxTime
                                                                    p25TxTime
#> [19] sdTimeToOutcome minTimeToOutcome p10TimeToOutcome
                                                                   p25TimeToOutcome
#> [28] irrLb95
                          irrUb95
                                               logRr
                                                                    seLogRr
#> <0 rows> (or 0-length row.names)
```

numOutcomes.

medianTxTim

medianTimeT

Analysis result objects are saved to the output folder and are aggregated by the summarizeAnalyses function which returns a results dataframe with one row for each result.

5 Correcting bias with EmpiricalCalibration

Effect estimates generated from observational data likely suffer from significant systematic bias. For example, this study design has the flaw that an exposure may look protective due to confounding by indication where patients are commonly exposed to a medication that is used to treat a comorbidity. For this reason, the SelfControlledCohort is often used with the EmpiricalCalibration package to adjust effect estimates using negative control cohorts.

See MethodEvaulation For more details on evaluating algorithms for population-level effect estimation in observational studies.