# Package 'CohortExplorer'

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Type Package	
Title An R package with a Shiny viewer to explore profiles of patients in a cohort	
Version 0.0.14	
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<b>Description</b> An R package with a Shiny viewer to explore profiles of patients in a cohort.	
<b>Depends</b> DatabaseConnector (>= 5.0.0),	
R (>= 4.0.0)	
Imports checkmate,	
clock,	
dplyr,	
lifecycle,	
ParallelLogger,	
rlang, stats	
Suggests rmarkdown, testthat,	
knitr,	
withr,	
SqlRender	
License Apache License	
RoxygenNote 7.2.3	
VignetteBuilder knitr	
Roxygen list(markdown = TRUE)	
Encoding UTF-8	
Language en-US	
<pre>URL https://ohdsi.github.io/CohortExplorer/, https:    //github.com/OHDSI/CohortExplorer</pre>	
<pre>BugReports https://github.com/OHDSI/CohortExplorer/issues</pre>	
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createCohortExplorerApp

Create Cohort explorer shiny app with person level data

#### **Description**

Export person level data from omop cdm tables for eligible persons in the cohort. Creates a folder with files that are part of the Cohort Explorer Shiny app. This app may then be run to review person level profiles.

### Usage

```
createCohortExplorerApp(
  connectionDetails = NULL,
  connection = NULL,
  cohortDatabaseSchema = NULL,
  cdmDatabaseSchema,
  vocabularyDatabaseSchema = cdmDatabaseSchema,
  tempEmulationSchema = getOption("sqlRenderTempEmulationSchema"),
  cohortTable = "cohort",
  cohortDefinitionId,
  cohortName = NULL,
  doNotExportCohortData = FALSE,
  sampleSize = 25,
  personIds = NULL,
  exportFolder,
  databaseId,
  shiftDates = FALSE,
  assignNewId = FALSE
)
```

#### **Arguments**

connectionDetails

An object of type connectionDetails as created using the createConnectionDetails function in the DatabaseConnector package. Can be left NULL if connection is provided.

connection

An object of type connection as created using the connect function in the DatabaseConnector package. Can be left NULL if connectionDetails is provided, in which case a new connection will be opened at the start of the function, and closed when the function finishes.

cohortDatabaseSchema

Schema name where your cohort tables reside. Note that for SQL Server, this should include both the database and schema name, for example 'scratch.dbo'.

cdmDatabaseSchema

Schema name where your omop cdm tables with person level data reside. Note that for SQL Server, this should include both the database and schema name, for example 'scratch.dbo'.

#### vocabularyDatabaseSchema

Schema name where your omop vocabulary tables reside. It maybe the cd-mDatabaseSchema. Note that for SQL Server, this should include both the database and schema name, for example 'scratch.dbo'.

#### tempEmulationSchema

Some database platforms like Oracle and Impala do not truly support temp tables. To emulate temp tables, provide a schema with write privileges where temp tables can be created.

cohortTable The name of the cohort table.

cohortDefinitionId

The cohort id to extract records.

cohortName (optional) Cohort Name

doNotExportCohortData

(Optional) Do you want to not export cohort data? If set to true, parameters cohortDefinitionId, cohort, cohortDatabaseSchema, cohortName will be ignored. The persons entire observation period would be considered the cohort. Cohort Name will be 'Observation Period', cohort id will be set to 0.

sampleSize (Optional, default = 20) The number of persons to randomly sample. Ignored, if

personId is given.

personIds (Optional) An array of personId's to look for in Cohort table and CDM.

exportFolder The folder where the output will be exported to. If this folder does not exist it

will be created.

databaseId A short string for identifying the database (e.g. 'Synpuf'). This will be displayed

in shiny app to toggle between databases. Should not have space or underscore

(\_).

shiftDates (Default = FALSE) Do you want to shift dates? This will help further de-

identify data. The shift is the process of recalibrating dates such that all persons

min(observation\_period\_start\_date) is 2000-01-01.

assignNewId (Default = FALSE) Do you want to assign a newId for persons. This will replace

the personId in the source with a randomly assigned newId.

#### **Examples**

```
## Not run:
connectionDetails <- createConnectionDetails(
  dbms = "postgresq1",
  server = "ohdsi.com",
  port = 5432,
  user = "me",
  password = "secure"
)

createCohortExplorerApp(
  connectionDetails = connectionDetails,
    cohortDefinitionId = 1234
)

## End(Not run)</pre>
```

 ${\tt exportCohortExplorerAppFiles}$ 

Copy shiny app files

## Description

Copy shiny app files.

## Usage

```
exportCohortExplorerAppFiles(exportFolder)
```

## Arguments

 ${\tt exportFolder}$ 

The folder where the output will be exported to. If this folder does not exist it will be created.

## **Examples**

```
## Not run:
exportCohortExplorerAppFiles(
   exportFolder = "output"
)
## End(Not run)
```

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```