# Package 'CDMConnector'

July 16, 2024

Title Connect to an OMOP Common Data Model

Version 1.5.0

**Description** Provides tools for working with observational health data in the Observational Medical Outcomes Partnership (OMOP) Common Data Model format with a pipe friendly syntax.

Common data model database table references are stored in a single compound object along with metadata.

**License** Apache License (>= 2)

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 https://github.com/darwin-eu/CDMConnector

BugReports https://github.com/darwin-eu/CDMConnector/issues

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'reexports-omopgenerics.R' 'generateCohortSet.R' 'generateConceptCohortSet.R' 'summariseQuantile.R' 'utils.R' 'validate.R' 'zzz-deprecated.R'
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appe	ndPermanent Run a dplyr query and add the result set to an existing	

# Description

Run a dplyr query and add the result set to an existing

# Usage

```
appendPermanent(x, name, schema = NULL)
append_permanent(x, name, schema = NULL)
```

# Arguments

X	A dplyr query
name	Name of the table to be appended. If it does not already exist it will be created.
schema	Schema where the table exists. Can be a length 1 or 2 vector. (e.g. schema = $"my\_schema"$ , schema = $c("my\_schema", "dbo"))$

# Value

A dplyr reference to the newly created table

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#### **Examples**

```
## Not run:
library(CDMConnector)
con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomia_dir())</pre>
concept <- dplyr::tbl(con, "concept")</pre>
# create a table
rxnorm_count <- concept %>%
  dplyr::filter(domain_id == "Drug") %>%
  dplyr::mutate(isRxnorm = (vocabulary_id == "RxNorm")) %>%
  dplyr::count(domain_id, isRxnorm) %>%
  compute("rxnorm_count")
# append to an existing table
rxnorm_count <- concept %>%
  dplyr::filter(domain_id == "Procedure") %>%
  dplyr::mutate(isRxnorm = (vocabulary_id == "RxNorm")) %>%
  dplyr::count(domain_id, isRxnorm) %>%
  appendPermanent("rxnorm_count")
DBI::dbDisconnect(con, shutdown = TRUE)
## End(Not run)
```

asDate

as.Date dbplyr translation wrapper

# **Description**

This is a workaround for using as.Date inside dplyr verbs against a database backend. This function should only be used inside dplyr verbs where the first argument is a database table reference. asDate must be unquoted with !! inside dplyr verbs (see example).

# Usage

```
asDate(x)
as_date(x)
```

#### **Arguments**

x an R expression

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#### **Examples**

assert\_tables

Assert that tables exist in a cdm object

# **Description**

A cdm object is a list of references to a subset of tables in the OMOP Common Data Model. If you write a function that accepts a cdm object as a parameter assert\_tables/assertTables will help you check that the tables you need are in the cdm object, have the correct columns/fields, and (optionally) are not empty.

# Usage

```
assert_tables(cdm, tables, empty.ok = FALSE, add = NULL)
assertTables(cdm, tables, empty.ok = FALSE, add = NULL)
```

### **Arguments**

cdm	A cdm object
tables	A character vector of table names to check.
empty.ok	Should an empty table (0 rows) be considered an error? TRUE or FALSE (default)
add	An optional AssertCollection created by checkmate::makeAssertCollection() that errors should be added to.

#### Value

Invisibly returns the cdm object

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#### **Examples**

```
## Not run:
# Use assertTables inside a function to check that tables exist
countDrugsByGender <- function(cdm) {
    assertTables(cdm, tables = c("person", "drug_era"), empty.ok = FALSE)

    cdm$person %>%
        dplyr::inner_join(cdm$drug_era, by = "person_id") %>%
        dplyr::count(.data$gender_concept_id, .data$drug_concept_id) %>%
        dplyr::collect()
}

library(CDMConnector)
con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomia_dir())
cdm <- cdm_from_con(con)

countDrugsByGender(cdm)

DBI::dbDisconnect(con, shutdown = TRUE)

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

assert\_write\_schema

Assert that cdm has a writable schema

#### **Description**

A cdm object can optionally contain a single schema in a database with write access. assert\_write\_schema checks that the cdm contains the "write\_schema" attribute and tests that local dataframes can be written to tables in this schema.

### Usage

```
assert_write_schema(cdm, add = NULL)
assertWriteSchema(cdm, add = NULL)
```

# Arguments

cdm A cdm object

add An optional AssertCollection created by checkmate::makeAssertCollection()

that errors should be added to.

#### Value

Invisibly returns the cdm object

benchmarkCDMConnector

benchmarkCDMConnector Run benchmark of tasks using CDMConnector

# **Description**

Run benchmark of tasks using CDMConnector

# Usage

```
benchmarkCDMConnector(cdm)
```

# **Arguments**

cdm

A CDM reference object

#### Value

a tibble with time taken for different analyses

# **Examples**

```
## Not run:
library(CDMConnector)
con <- DBI::dbConnect(duckdb::duckdb(), eunomia_dir())
cdm <- cdm_from_con(con, cdm_schema = "main", write_schema = "main")
benchmarkCDMConnector(cdm)

DBI::dbDisconnect(con, shutdown = TRUE)
## End(Not run)</pre>
```

cdmCon

Get underlying database connection

# **Description**

Get underlying database connection

# Usage

```
cdmCon(cdm)
```

# **Arguments**

cdm

A cdm reference object created by cdm\_from\_con

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#### Value

A reference to the database containing tables in the cdm reference

# **Examples**

cdmDisconnect

Disconnect the connection of the cdm object

# Description

This function will disconnect from the database as well as drop "temporary" tables that were created on database systems that do not support actual temporary tables. Currently temp tables are emulated on Spark/Databricks systems.

# Usage

```
cdmDisconnect(cdm)
cdm_disconnect(cdm)
```

# **Arguments**

cdm

cdm reference

cdmFlatten

Flatten a cdm into a single observation table

# Description

This experimental function transforms the OMOP CDM into a single observation table. This is only recommended for use with a filtered CDM or a cdm that is small in size.

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### Usage

```
cdmFlatten(
  cdm,
  domain = c("condition", "drug", "procedure"),
  includeConceptName = TRUE
)

cdm_flatten(
  cdm,
  domain = c("condition", "drug", "procedure"),
  include_concept_name = TRUE
)
```

# **Arguments**

Should concept\_name and type\_concept\_name be include in the output table? TRUE (default) or FALSE

#### **Details**

### [Experimental]

#### Value

A lazy query that when evaluated will result in a single cdm table

```
## Not run:
library(CDMConnector)
library(dplyr, warn.conflicts = FALSE)
con <- DBI::dbConnect(duckdb::duckdb(), eunomia_dir())</pre>
cdm <- cdm_from_con(con, cdm_schema = "main")</pre>
all_observations <- cdmSubset(cdm, personId = c(2, 18, 42)) %>%
  cdmFlatten() %>%
  collect()
all_observations
#> # A tibble: 213 × 8
    person_id observation_. start_date end_date type_. domain obser. type_.
#>
     <dbl> <dbl> <date> <date>
                                                        <dbl> <chr> <chr>
                                                                               <chr>
          2 40213201 1986-09-09 1986-09-09 5.81e5 drug pneumo <NA>
18 4116491 1997-11-09 1998-01-09 3.20e4 condi Escher <NA>
#> 1
#> 2
```

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```
18
                     40213227 2017-01-04 2017-01-04 5.81e5 drug
                                                                  tetanu
#>
                     4156265 1974-06-13 1974-06-27 3.20e4 condi
            42
                                                                 Facial
#>
  5
            18
                     40213160 1966-02-23 1966-02-23 5.81e5 drug
                                                                         <NA>
                                                                 poliov
#> 6
            42
                     4198190 1933-10-29 1933-10-29 3.80e7 proce
                                                                 Append
                                                                         <NA>
#> 7
            2
                      4109685 1952-07-13 1952-07-27 3.20e4 condi Lacera <NA>
#> 8
            18
                     40213260 2017-01-04 2017-01-04 5.81e5 drug
                                                                  zoster <NA>
#> 9
            42
                      4151422 1985-02-03 1985-02-03 3.80e7 proce Sputum <NA>
#> 10
                      4163872 1993-03-29 1993-03-29 3.80e7 proce Plain
#> # ... with 203 more rows, and abbreviated variable names observation_concept_id,
      type_concept_id, observation_concept_name, type_concept_name
DBI::dbDisconnect(con, shutdown = TRUE)
## End(Not run)
```

cdmName

Get the CDM name

### **Description**

Extract the CDM name attribute from a cdm\_reference object

#### Usage

```
cdmName(cdm)
cdm_name(cdm)
```

### **Arguments**

cdm

A cdm object

### Value

The name of the CDM as a character string

```
## Not run:
library(CDMConnector)
con <- DBI::dbConnect(duckdb::duckdb(), eunomia_dir())
cdm <- cdm_from_con(con, cdm_schema = "main", write_schema = "main")
cdmName(cdm)
#> [1] "eunomia"

DBI::dbDisconnect(con, shutdown = TRUE)
## End(Not run)
```

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cdmSample

Subset a cdm object to a random sample of individuals

#### **Description**

cdmSample takes a cdm object and returns a new cdm that includes only a random sample of persons in the cdm. Only person\_ids in both the person table and observation\_period table will be considered.

# Usage

```
cdmSample(cdm, n, seed = sample.int(1e+06, 1), name = "person_sample")
cdm_sample(cdm, n, seed = sample.int(1e+06, 1), name = "person_sample")
```

#### **Arguments**

cdm A cdm\_reference object.

Number of persons to include in the cdm.seedSeed for the random number generator.

name Name of the table that will contain the sample of persons.

#### **Details**

# [Experimental]

#### Value

A modified cdm\_reference object where all clinical tables are lazy queries pointing to subset

```
## Not run:
library(CDMConnector)
library(dplyr, warn.conflicts = FALSE)

con <- DBI::dbConnect(duckdb::duckdb(), eunomia_dir())

cdm <- cdm_from_con(con, cdm_schema = "main")

cdmSampled <- cdmSample(cdm, n = 2)

cdmSampled$person %>%
    select(person_id)

#> # Source: SQL [2 x 1]

#> # Database: DuckDB 0.6.1

#> person_id

#> <dbl>
```

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```
#> 1    155
#> 2    3422

DBI::dbDisconnect(con, shutdown = TRUE)
## End(Not run)
```

cdmSubset

Subset a cdm object to a set of persons

# **Description**

cdmSubset takes a cdm object and a list of person IDs as input. It returns a new cdm that includes data only for persons matching the provided person IDs. Generated cohorts in the cdm will also be subset to the IDs provided.

# Usage

```
cdmSubset(cdm, personId)
cdm_subset(cdm, person_id)
```

### **Arguments**

```
cdm A cdm_reference object person_id, personId
```

A numeric vector of person IDs to include in the cdm

#### **Details**

# [Experimental]

#### Value

A modified cdm\_reference object where all clinical tables are lazy queries pointing to subset

```
## Not run:
library(CDMConnector)
library(dplyr, warn.conflicts = FALSE)

con <- DBI::dbConnect(duckdb::duckdb(), eunomia_dir())

cdm <- cdm_from_con(con, cdm_schema = "main")

cdm2 <- cdmSubset(cdm, personId = c(2, 18, 42))

cdm2$person %>%
```

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```
select(1:3)
#> # Source:
               SQL [3 x 3]
#> # Database: DuckDB 0.6.1
     person_id gender_concept_id year_of_birth
#>
         <dbl>
                           <dbl>
                                          <dbl>
#> 1
            2
                             8532
                                           1920
#> 2
            18
                             8532
                                           1965
            42
                             8532
                                           1909
DBI::dbDisconnect(con, shutdown = TRUE)
## End(Not run)
```

cdmSubsetCohort

Subset a cdm to the individuals in one or more cohorts

### **Description**

cdmSubset will return a new cdm object that contains lazy queries pointing to each of the cdm tables but subset to individuals in a generated cohort. Since the cdm tables are lazy queries, the subset operation will only be done when the tables are used. computeQuery can be used to run the SQL used to subset a cdm table and store it as a new table in the database.

# Usage

```
cdmSubsetCohort(cdm, cohortTable = "cohort", cohortId = NULL, verbose = FALSE)

cdm_subset_cohort(
    cdm,
    cohort_table = "cohort",
    cohort_id = NULL,
    verbose = FALSE
)
```

# **Arguments**

```
cdm A cdm_reference object

cohortTable, cohort_table

The name of a cohort table in the cdm reference

cohortId, cohort_id

IDs of the cohorts that we want to subset from the cohort table. If NULL (default) all cohorts in cohort table are considered.

verbose

Should subset messages be printed? TRUE or FALSE (default)
```

#### **Details**

# [Experimental]

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#### Value

A modified cdm\_reference with all clinical tables subset to just the persons in the selected cohorts.

### **Examples**

```
## Not run:
library(CDMConnector)
library(dplyr, warn.conflicts = FALSE)
con <- DBI::dbConnect(duckdb::duckdb(), eunomia_dir())</pre>
cdm <- cdm_from_con(con, cdm_schema = "main", write_schema = "main")</pre>
# generate a cohort
path <- system.file("cohorts2", mustWork = TRUE, package = "CDMConnector")</pre>
cohortSet <- readCohortSet(path) %>%
  filter(cohort_name == "GIBleed_male")
# subset cdm to persons in the generated cohort
cdm <- generateCohortSet(cdm, cohortSet = cohortSet, name = "gibleed")</pre>
cdmGiBleed <- cdmSubsetCohort(cdm, cohortTable = "gibleed")</pre>
cdmGiBleed$person %>%
  tally()
#> # Source: SQL [1 x 1]
#> # Database: DuckDB 0.6.1
#>
    <dbl>
#> 1 237
cdm$person %>%
  tally()
#> # Source: SQL [1 x 1]
#> # Database: DuckDB 0.6.1
#>
         n
#>
    <dbl>
#> 1 2694
DBI::dbDisconnect(con, shutdown = TRUE)
## End(Not run)
```

cdmWriteSchema

Get cdm write schema

#### **Description**

Get cdm write schema

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# Usage

```
cdmWriteSchema(cdm)
```

# **Arguments**

cdm

A cdm reference object created by cdm\_from\_con

#### Value

The database write schema

# **Examples**

 $cdm\_from\_con$ 

Create a CDM reference object from a database connection

# **Description**

Create a CDM reference object from a database connection

# Usage

```
cdm_from_con(
   con,
   cdm_schema,
   write_schema,
   cohort_tables = NULL,
   cdm_version = "5.3",
   cdm_name = NULL,
   achilles_schema = NULL,
   .soft_validation = FALSE,
   write_prefix = NULL
)

cdmFromCon(
   con,
```

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```
cdmSchema,
writeSchema,
cohortTables = NULL,
cdmVersion = "5.3",
cdmName = NULL,
achillesSchema = NULL,
.softValidation = FALSE,
writePrefix = NULL
```

# Arguments

con

A DBI database connection to a database where an OMOP CDM v5.4 or v5.3 instance is located.

cdm\_schema, cdmSchema

The schema where the OMOP CDM tables are located. Defaults to NULL.

write\_schema, writeSchema

An optional schema in the CDM database that the user has write access to.

cohort\_tables, cohortTables

A character vector listing the cohort table names to be included in the CDM object.

cdm\_version, cdmVersion

The version of the OMOP CDM: "5.3" (default), "5.4", "auto". "auto" attempts to automatically determine the cdm version using heuristics. Cohort tables must be in the write\_schema.

cdm\_name, cdmName

The name of the CDM. If NULL (default) the cdm\_source\_name . field in the CDM\_SOURCE table will be used.

achilles\_schema, achillesSchema

An optional schema in the CDM database that contains achilles tables.

.soft\_validation, .softValidation

Normally the observation period table should not have overlapping observation periods for a single person. If .softValidation is TRUE the validation check that looks for overlapping observation periods will be skipped. Other analytic packages may break or produce incorrect results if softValidation is TRUE and the observation period table contains overlapping observation periods.

write\_prefix, writePrefix

A prefix that will be added to all tables created in the write\_schema. This can be used to create namespace in your database write\_schema for your tables.

#### **Details**

cdm\_from\_con / cdmFromCon creates a new cdm reference object from a DBI compliant database connection. In addition to the connection the user needs to pass in the schema in the database where the cdm data can be found as well as another schema where the user has write access to create tables. Nearly all downstream analytic packages need the ability to create temporary data in the database so the write\_schema is required.

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Some database systems have the idea of a catalog or a compound schema with two components. See examples below for how to pass in catalogs and schemas.

You can also specify a write\_prefix. This is a short character string that will be added to any tables created in the write\_schema effectively a namespace in the schema just for your analysis. This makes it easy to ensure you do not overwrite someone elses tables if the write\_schema is shared and allows you to easily clean up tables by dropping all tables that start with the prefix. The prefix is considered part of the write\_schema since it is effectively a sub-schema. See examples.

#### Value

A list of dplyr database table references pointing to CDM tables

```
## Not run:
library(CDMConnector)
con <- DBI::dbConnect(duckdb::duckdb(), eunomia_dir())</pre>
# minimal example
cdm <- cdm_from_con(con,</pre>
                     cdm_schema = "main",
                     write_schema = "scratch")
cdm <- cdm_from_con(con,</pre>
                     cdm_schema = "main",
                     write_schema = "scratch",
                     write_prefix = "tmp_")
# There are a few differen options for using catalogs
cdm <- cdm_from_con(con,</pre>
                     cdm_schema = "catalog.main",
                     write_schema = "catalog.scratch",
                     write_prefix = "tmp_")
cdm <- cdm_from_con(con,</pre>
                     cdm_schema = c(catalog = "catalog", schema = "main"),
                     write_schema = c(catalog = "catalog", schema = "scratch"))
cdm <- cdm_from_con(con,</pre>
                     cdm_schema = c("catalog", "main"),
                     write_schema = c("catalog", "scratch"))
cdm <- cdm_from_con(con,</pre>
                     cdm_schema = c(catalog = "catalog", schema = "main"),
                     write_schema = c(catalog = "catalog",
                                       schema = "scratch",
                                       prefix = "tmp_"))
 DBI::dbDisconnect(con)
## End(Not run)
```

cdm\_from\_environment Create a CDM object from a pre-defined set of environment variables

# **Description**

This function is intended to be used with the Darwin execution engine. The execution engine runs OHDSI studies in a pre-defined runtime environment and makes several environment variables available for connecting to a CDM database. Programmer writing code to run on the execution engine and simply use cdm <- cdm\_from\_environment() to create a cdm reference object to use for their analysis and the database connection and cdm object should be automatically created. This obviates the need for site specific code for connecting to the database and creating the cdm reference object.

### Usage

```
cdm_from_environment(write_prefix = "")
```

#### **Arguments**

write\_prefix (string) An optional prefix to use for all tables written to the CDM.

### **Details**

The environment variables used by this function and provided by the execution engine are listed below.

- DBMS\_TYPE: one of "postgresql", "sql server", "redshift", "duckdb", "snowflake".
- DATA\_SOURCE\_NAME: a free text name for the CDM given by the person running the study.
- CDM\_VERSION: one of "5.3", "5.4".
- DBMS\_CATALOG: The database catalog. Important primarily for compound schema names used in SQL Server and Snowflake.
- DBMS\_SERVER: The database server URL.
- DBMS\_NAME: The database name used for creating the connection.
- DBMS\_PORT: The database port number.
- DBMS\_USERNAME: The database username needed to authenticate.
- DBMS\_PASSWORD: The database password needed to authenticate.
- CDM\_SCHEMA: The schema name where the OMOP CDM is located in the database.
- WRITE\_SCHEMA: The shema where the user has write access and tables will be created during study execution.

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# Value

A cdm\_reference object

#### **Examples**

```
## Not run:
library(CDMConnector)

# This will only work in an evironment where the proper variables are present.
cdm <- cdm_from_environment()

# Proceed with analysis using the cdm object.

# Close the database connection when done.
cdm_disconnect(cdm)

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

cdm\_from\_files

Create a CDM reference from a folder containing parquet, csv, or feather files

# **Description**

Create a CDM reference from a folder containing parquet, csv, or feather files

# Usage

```
cdm_from_files(
  path,
  format = "auto",
  cdm_version = "5.3",
  cdm_name = NULL,
  as_data_frame = TRUE
)

cdmFromFiles(
  path,
  format = "auto",
  cdmVersion = "5.3",
  cdmName = NULL,
  asDataFrame = TRUE
)
```

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### **Arguments**

path A folder where an OMOP CDM v5.4 instance is located.

format What is the file format to be read in? Must be "auto" (default), "parquet", "csv",

"feather".

cdm\_version, cdmVersion

The version of the cdm (5.3 or 5.4)

cdm\_name, cdmName

A name to use for the cdm.

as\_data\_frame, asDataFrame

TRUE (default) will read files into R as dataframes. FALSE will read files into

R as Arrow Datasets.

#### Value

A list of dplyr database table references pointing to CDM tables

cdm\_from\_tables

Create a cdm object from local tables

# Description

Create a cdm object from local tables

#### Usage

```
cdm_from_tables(tables, cdm_name, cohort_tables = list(), cdm_version = NULL)
```

# **Arguments**

tables List of tables to be part of the cdm object.

cdm\_name Name of the cdm object.

cohort\_tables List of tables that contains cohort, cohort\_set and cohort\_attrition can be pro-

vided as attributes.

cdm\_version Version of the cdm\_reference

#### Value

A cdm\_reference object.

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#### **Examples**

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)

person <- tibble(
    person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
    race_concept_id = 0, ethnicity_concept_id = 0
)
observation_period <- tibble(
    observation_period_id = 1, person_id = 1,
    observation_period_start_date = as.Date("2000-01-01"),
    observation_period_end_date = as.Date("2025-12-31"),
    period_type_concept_id = 0
)
cdm <- cdm_from_tables(
    tables = list("person" = person, "observation_period" = observation_period),
    cdm_name = "test"
)</pre>
```

cdm\_select\_tbl

Select a subset of tables in a cdm reference object

# **Description**

This function uses syntax similar to dplyr::select and can be used to subset a cdm reference object to a specific tables

#### Usage

```
cdm_select_tbl(cdm, ...)
```

#### **Arguments**

cdm A cdm reference object created by cdm\_from\_con... One or more table names of the tables of the cdm object. tidyselect is supported, see dplyr::select() for details on the semantics.

#### Value

A cdm reference object containing the selected tables

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomia_dir())
cdm <- cdm_from_con(con, "main")</pre>
```

22 cohortSet

```
cdm_select_tbl(cdm, person)
cdm_select_tbl(cdm, person, observation_period)
cdm_select_tbl(cdm, tbl_group("vocab"))
cdm_select_tbl(cdm, "person")

DBI::dbDisconnect(con)

## End(Not run)
```

cohortAttrition

Get attrition table from a cohort\_table object

# Description

Get attrition table from a cohort\_table object

# Usage

```
cohortAttrition(x)
cohort_attrition(x)
```

# Arguments

Х

A cohort\_table object

cohortSet

Get cohort settings from a cohort\_table object

# Description

Get cohort settings from a cohort\_table object

# Usage

```
cohortSet(x)
cohort_set(x)
```

# **Arguments**

Х

A cohort\_table object

cohort\_count 23

 $cohort\_count$ 

Get cohort counts from a generated\_cohort\_set object.

# **Description**

Get cohort counts from a generated\_cohort\_set object.

# Usage

```
cohort_count(cohort)
```

### **Arguments**

cohort

A generated\_cohort\_set object.

#### Value

A table with the counts.

# **Examples**

```
## Not run:
library(CDMConnector)
library(dplyr)

con <- DBI::dbConnect(duckdb::duckdb(), eunomia_dir())
cdm <- cdm_from_con(con = con, cdm_schema = "main", write_schema = "main")
cdm <- generateConceptCohortSet(
   cdm = cdm, conceptSet = list(pharyngitis = 4112343), name = "new_cohort"
)
cohort_count(cdm$new_cohort)

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

cohort\_erafy

Collapse cohort records within a certain number of days

# Description

Collapse cohort records within a certain number of days

# Usage

```
cohort_erafy(x, gap)
cohortErafy(x, gap)
```

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#### **Arguments**

x A generated cohort set

gap When two cohort records are 'gap' days apart or less the periods will be col-

lapsed into a single record

#### Value

A lazy query on a generated cohort set

cohort\_union

Union all cohorts in a cohort set with cohorts in a second cohort set

### **Description**

Union all cohorts in a cohort set with cohorts in a second cohort set

# Usage

```
cohort_union(x, y)
cohortUnion(x, y)
```

#### **Arguments**

x A tbl reference to a cohort table with one or more generated cohorts

y A tbl reference to a cohort table with one generated cohort

#### Value

A lazy query that when executed will resolve to a new cohort table with one the same cohort\_definitions\_ids in x resulting from the union of all cohorts in x with the single cohort in y cohort table

computeQuery

Execute dplyr query and save result in remote database

# **Description**

This function is a wrapper around dplyr::compute that is tested on several database systems. It is needed to handle edge cases where dplyr::compute does not produce correct SQL.

computeQuery 25

# Usage

```
computeQuery(
  name = uniqueTableName(),
  temporary = TRUE,
  schema = NULL,
 overwrite = TRUE,
)
compute_query(
 х,
 name = uniqueTableName(),
  temporary = TRUE,
  schema = NULL,
 overwrite = TRUE,
)
```

### **Arguments**

Х A dplyr query

name The name of the table to create.

Should the table be temporary: TRUE (default) or FALSE temporary

The schema where the table should be created. Ignored if temporary = TRUE. schema overwrite

Should the table be overwritten if it already exists: TRUE (default) or FALSE

Ignored if temporary = TRUE.

Further arguments passed on the dplyr::compute . . .

#### Value

A dplyr::tbl() reference to the newly created table.

```
## Not run:
library(CDMConnector)
con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomia_dir())</pre>
cdm <- cdm_from_con(con, "main")</pre>
# create a temporary table in the remote database from a dplyr query
drugCount <- cdm$concept %>%
  dplyr::count(domain_id == "Drug") %>%
  computeQuery()
# create a permanent table in the remote database from a dplyr query
drugCount <- cdm$concept %>%
```

26 copy\_cdm\_to

```
dplyr::count(domain_id == "Drug") %>%
  computeQuery("tmp_table", temporary = FALSE, schema = "main")
DBI::dbDisconnect(con, shutdown = TRUE)
## End(Not run)
```

copy\_cdm\_to

Copy a cdm object from one database to another

# Description

It may be helpful to be able to easily copy a small test cdm from a local database to a remote for testing. copy\_cdm\_to takes a cdm object and a connection. It copies the cdm to the remote database connection. CDM tables can be prefixed in the new database allowing for multiple cdms in a single shared database schema.

### Usage

```
copy_cdm_to(con, cdm, schema, overwrite = FALSE)
copyCdmTo(con, cdm, schema, overwrite = FALSE)
```

### **Arguments**

con A DBI datbase connection created by DBI::dbConnect

schema schema name in the remote database where the user has write permission

overwrite Should the cohort table be overwritten if it already exists? TRUE or FALSE

(default)

# **Details**

### [Experimental]

#### Value

A cdm reference object pointing to the newly created cdm in the remote database

dateadd 27

dateadd

Add days or years to a date in a dplyr query

#### **Description**

This function must be "unquoted" using the "bang bang" operator (!!). See example.

# Usage

```
dateadd(date, number, interval = "day")
```

# **Arguments**

The name of a date column in the database table as a character string number

The number of units to add. Can be a positive or negative whole number.

interval The units to add. Must be either "day" (default) or "year"

#### Value

Platform specific SQL that can be used in a dplyr query.

#### **Examples**

datediff

Compute the difference between two days

# **Description**

This function must be "unquoted" using the "bang bang" operator (!!). See example.

#### Usage

```
datediff(start, end, interval = "day")
```

28 datepart

# **Arguments**

start The name of the start date column in the database as a string.

end The name of the end date column in the database as a string.

interval The units to use for difference calculation. Must be either "day" (default) or

"year".

#### Value

Platform specific SQL that can be used in a dplyr query.

#### **Examples**

datepart

Extract the day, month or year of a date in a dplyr pipeline

#### **Description**

Extract the day, month or year of a date in a dplyr pipeline

#### **Usage**

```
datepart(date, interval = "year", dbms = NULL)
```

# **Arguments**

date Character string that represents to a date column.

interval Interval to extract from a date. Valid options are "year", "month", or "day".

dbms Database system, if NULL it is auto detected.

dbms 29

#### **Examples**

dbms

Get the database management system (dbms) from a cdm\_reference or DBI connection

# **Description**

Get the database management system (dbms) from a cdm\_reference or DBI connection

### Usage

dbms(con)

# **Arguments**

con

A DBI connection or cdm\_reference

### Value

A character string representing the dbms that can be used with SqlRender

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomia_dir())
cdm <- cdm_from_con(con)
dbms(cdm)
dbms(con)
## End(Not run)</pre>
```

30 downloadEunomiaData

dbSource

Create a source for a cdm in a database.

# Description

Create a source for a cdm in a database.

# Usage

```
dbSource(con, writeSchema)
```

# Arguments

con Connection to a database.

writeSchema Schema where cohort tables are. You must have read and write access to it.

downloadEunomiaData

Download Eunomia data files

# **Description**

Download the Eunomia data files from https://github.com/darwin-eu/EunomiaDatasets

# Usage

```
downloadEunomiaData(
  datasetName = "GiBleed",
  cdmVersion = "5.3",
  pathToData = Sys.getenv("EUNOMIA_DATA_FOLDER"),
  overwrite = FALSE
)

download_eunomia_data(
  dataset_name = "GiBleed",
  cdm_version = "5.3",
  path_to_data = Sys.getenv("EUNOMIA_DATA_FOLDER"),
  overwrite = FALSE
)
```

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#### **Arguments**

overwrite Control whether the existing archive file will be overwritten should it already exist.

dataset\_name, datasetName

The data set name as found on https://github.com/darwin-eu/EunomiaDatasets. The data set name corresponds to the folder with the data set ZIP files

cdm\_version, cdmVersion

The OMOP CDM version. This version will appear in the suffix of the data file, for example: synpuf\_5.3.zip. Must be '5.3' (default) or '5.4'.

path\_to\_data, pathToData

The path where the Eunomia data is stored on the file system., By default the value of the environment variable "EUNOMIA\_DATA\_FOLDER" is used.

#### Value

Invisibly returns the destination if the download was successful.

# **Examples**

```
## Not run:
downloadEunomiaData("GiBleed")
## End(Not run)
```

eunomiaDir

Create a copy of an example OMOP CDM dataset

#### **Description**

Creates a copy of a Eunomia database, and returns the path to the new database file. If the dataset does not yet exist on the user's computer it will attempt to download the source data to the the path defined by the EUNOMIA\_DATA\_FOLDER environment variable.

#### Usage

```
eunomiaDir(
  datasetName = "GiBleed",
  cdmVersion = "5.3",
  databaseFile = tempfile(fileext = ".duckdb")
)

eunomia_dir(
  dataset_name = "GiBleed",
  cdm_version = "5.3",
  database_file = tempfile(fileext = ".duckdb")
)
```

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#### **Arguments**

```
datasetName, dataset_name
```

One of "GiBleed" (default), "synthea-allergies-10k", "synthea-anemia-10k", "synthea-breast\_cancer-10k", "synthea-contraceptives-10k", "synthea-covid19-10k", "synthea-covid19-200k", "synthea-dermatitis-10k", "synthea-heart-10k", "synthea-hiv-10k", "synthea-lung\_cancer-10k", "synthea-medications-10k", "synthea-metabolic\_syndrome-10k", "synthea-opioid\_addiction-10k", "synthea-rheumatoid\_arthritis-10k", "synthea-snf-10k", "synthea-surgery-10k", "synthea-total\_joint\_replacement-10k", "synthea-veteran\_prostate\_cancer-10k", "synthea-veterans-10k", "synthea-weight\_loss-10k"

cdmVersion, cdm\_version

The OMOP CDM version. Must be "5.3" or "5.4". databaseFile, database\_file  $\,$ 

The full path to the new copy of the example CDM dataset.

#### Value

The file path to the new Eunomia dataset copy

### **Examples**

```
## Not run:
  con <- DBI::dbConnect(duckdb::duckdb(), eunomiaDir("GiBleed"))
  DBI::dbDisconnect(con, shutdown = TRUE)
## End(Not run)</pre>
```

#### Description

Has the Eunomia dataset been cached?

#### Usage

```
eunomia_is_available(dataset_name = "GiBleed", cdm_version = "5.3")
eunomiaIsAvailable(datasetName = "GiBleed", cdmVersion = "5.3")
```

### **Arguments**

```
{\bf Name\ of\ the\ Eunomia\ dataset\ to\ check.\ Defaults\ to\ "GiBleed".} {\bf cdm\_version,\ cdmVersion}
```

Version of the Eunomia dataset to check. Must be "5.3" or "5.4".

#### Value

TRUE if the eunomia example dataset is available and FASLE otherwise

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exampleDatasets

List the available example CDM datasets

#### **Description**

List the available example CDM datasets

#### Usage

```
exampleDatasets()
example_datasets()
```

#### Value

A character vector with example CDM dataset identifiers

# **Examples**

```
## Not run:
library(CDMConnector)
exampleDatasets()[1]
#> [1] "GiBleed"

con <- DBI::dbConnect(duckdb::duckdb(), eunomiaDir("GiBleed"))
cdm <- cdm_from_con(con)

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

generateCohortSet

Generate a cohort set on a cdm object

#### **Description**

A "chort\_table" object consists of four components

- A remote table reference to an OHDSI cohort table with at least the columns: cohort\_definition\_id, subject\_id, cohort\_start\_date, cohort\_end\_date. Additional columns are optional and some analytic packages define additional columns specific to certain analytic cohorts.
- A **settings attribute** which points to a remote table containing cohort settings including the names of the cohorts.
- An **attrition attribute** which points to a remote table with attrition information recorded during generation. This attribute is optional. Since calculating attrition takes additional compute it can be skipped resulting in a NULL attrition attribute.
- A cohortCounts attribute which points to a remote table containing cohort counts

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Each of the three attributes are tidy tables. The implementation of this object is experimental and user feedback is welcome.

#### [Experimental]

One key design principle is that cohort\_table objects are created once and can persist across analysis execution but should not be modified after creation. While it is possible to modify a cohort\_table object doing so will invalidate it and it's attributes may no longer be accurate.

# Usage

```
generateCohortSet(
  cdm,
  cohortSet,
  name,
  computeAttrition = TRUE,
  overwrite = TRUE
)

generate_cohort_set(
  cdm,
  cohort_set,
  name = "cohort",
  compute_attrition = TRUE,
  overwrite = TRUE
)
```

#### **Arguments**

A cdm reference created by CDMConnector. write\_schema must be specified.

Name of the cohort table to be created. This will also be used as a prefix for the cohort attribute tables.

overwrite Should the cohort table be overwritten if it already exists? TRUE (default) or FALSE

cohort\_set, cohortSet

Can be a cohortSet object created with readCohortSet()

compute\_attrition, computeAttrition

Should attrition be computed? TRUE (default) or FALSE

```
print(cdm$cohort)
attrition(cdm$cohort)
settings(cdm$cohort)
cohortCount(cdm$cohort)
## End(Not run)
```

generateConceptCohortSet

Create a new generated cohort set from a list of concept sets

### **Description**

Generate a new cohort set from one or more concept sets. Each concept set will result in one cohort and represent the time during which the concept was observed for each subject/person. Concept sets can be passed to this function as:

- A named list of numeric vectors, one vector per concept set
- A named list of Capr concept sets

Clinical observation records will be looked up in the respective domain tables using the vocabulary in the CDM. If a required domain table does not exist in the cdm object a warning will be given. Concepts that are not in the vocabulary or in the data will be silently ignored. If end dates are missing or do not exist, as in the case of the procedure and observation domains, the the start date will be used as the end date.

#### Usage

```
generateConceptCohortSet(
  cdm.
  conceptSet = NULL,
  name,
  limit = "first",
  requiredObservation = c(0, 0),
  end = "observation_period_end_date",
  subsetCohort = NULL,
  subsetCohortId = NULL,
  overwrite = TRUE
)
generate_concept_cohort_set(
  cdm,
  concept_set = NULL,
  name = "cohort",
  limit = "first",
  required_observation = c(0, 0),
```

```
end = "observation_period_end_date",
subset_cohort = NULL,
subset_cohort_id = NULL,
overwrite = TRUE
)
```

#### **Arguments**

 $\label{lem:composition} A \ cdm \ reference \ object \ created \ by \ CDMConnector:: cdmFromCon \ or \ CDMConnector:: cdm_from\_con \ conceptSet, \ concept\_set$ 

A named list of numeric vectors or a Concept Set Expression created omopgenerics::newConceptSetExp

name

The name of the new generated cohort table as a character string

limit

Include "first" (default) or "all" occurrences of events in the cohort

- "first" will include only the first occurrence of any event in the concept set in the cohort.
- "all" will include all occurrences of the events defined by the concept set in the cohort.

requiredObservation, required\_observation

A numeric vector of length 2 that specifies the number of days of required observation time prior to index and post index for an event to be included in the cohort.

end

How should the cohort\_end\_date be defined?

- "observation\_period\_end\_date" (default): The earliest observation\_period\_end\_date after the event start date
- numeric scalar: A fixed number of days from the event start date
- "event\_end\_date": The event end date. If the event end date is not populated then the event start date will be used

 $subsetCohort, subset\_cohort$ 

A cohort table containing the individuals for which to generate cohorts for. Only individuals in the cohort table will appear in the created generated cohort set.

subsetCohortId, subset\_cohort\_id

A set of cohort IDs from the cohort table for which to include. If none are provided, all cohorts in the cohort table will be included.

overwrite

Should the cohort table be overwritten if it already exists? TRUE (default) or FALSE.

#### Value

A cdm reference object with the new generated cohort set table added

inSchema 37

inSchema	Helper for working with compound schemas
----------	--

## **Description**

This is similar to dbplyr::in\_schema but has been tested across multiple database platforms. It only exists to work around some of the limitations of dbplyr::in\_schema.

#### Usage

```
inSchema(schema, table, dbms = NULL)
in_schema(schema, table, dbms = NULL)
```

## Arguments

schema A schema name as a character string
table A table name as character string

dbms The name of the database management system as returned by dbms (connection)

#### Value

A DBI::Id that represents a qualified table and schema

# **Description**

Intersect all cohorts in a single cohort table

## Usage

```
intersect_cohorts(x, cohort_definition_id = 1L)
intersectCohorts(x, cohort_definition_id = 1L)
```

## **Arguments**

## Value

A lazy query that when executed will resolve to a new cohort table with one cohort\_definition\_id resulting from the intersection of all cohorts in the original cohort table

list\_tables

List tables in a schema

## **Description**

DBI::dbListTables can be used to get all tables in a database but not always in a specific schema. listTables will list tables in a schema.

# Usage

```
list_tables(con, schema = NULL)
listTables(con, schema = NULL)
```

## Arguments

con A DBI connection to a database

schema The name of a schema in a database. If NULL, returns DBI::dbListTables(con).

## Value

A character vector of table names

## **Examples**

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomia_dir())
listTables(con, schema = "main")
## End(Not run)</pre>
```

```
new_generated_cohort_set
```

Constructor for cohort\_table objects

## **Description**

## [Superseded]

#### **Usage**

```
new_generated_cohort_set(
  cohort_ref,
  cohort_set_ref = NULL,
  cohort_attrition_ref = NULL,
  cohort_count_ref = NULL,
  overwrite
)

newGeneratedCohortSet(
  cohortRef,
  cohortSetRef = NULL,
  cohortAttritionRef = NULL,
  cohortCountRef = NULL,
  overwrite
)
```

## Arguments

cohort\_ref, cohortRef

A tbl\_sql object that points to a remote cohort table with the following first four columns: cohort\_definition\_id, subject\_id, cohort\_start\_date, cohort\_end\_date. Additional columns are optional.

cohort\_set\_ref, cohortSetRef

A tbl\_sql object that points to a remote table with the following first two columns: cohort\_definition\_id, cohort\_name. Additional columns are optional. cohort\_definition\_id should be a primary key on this table and uniquely identify rows.

cohort\_attrition\_ref, cohortAttritionRef

A tbl\_sql object that points to an attrition table in a remote database with the first column being cohort\_definition\_id.

cohort\_count\_ref, cohortCountRef

A tbl\_sql object that points to a cohort\_count table in a remote database with columns cohort\_definition\_id, cohort\_entries, cohort\_subjects.

overwrite

Should tables be overwritten if they already exist? TRUE or FALSE (default)

#### **Details**

Please use omopgenerics::newCohortTable() instead.

This constructor function is to be used by analytic package developers to create cohort\_table objects.

A cohort\_table is a set of person-time from an OMOP CDM database. A cohort\_table can be represented by a table with three columns: subject\_id, cohort\_start\_date, cohort\_end\_date. Subject\_id is the same as person\_id in the OMOP CDM. A cohort\_table is a collection of one or more cohort\_table and can be represented as a table with four columns: cohort\_definition\_id, subject\_id, cohort\_start\_date, cohort\_end\_date.

This constructor function defines the cohort\_table object in R.

The object is an extension of a tbl\_sql object defined in dplyr. This is a lazy database query that points to a cohort table in the database with at least the columns cohort\_definition\_id, subject\_id, cohort\_start\_date, cohort\_end\_date. The table could optionally have more columns as well.

In addition the cohort\_table object has three optional attributes. These are: cohort\_set, cohort\_attrition, cohort\_count. Each of these attributes is also a lazy SQL query (tbl\_sql) that points to a table in a database and is described below.

#### cohort set:

cohort\_set is a table with one row per cohort\_definition\_id. The first two columns of the cohort\_set table are: cohort\_definition\_id, and cohort\_name. Additional columns can be added. The cohort\_set table is meant to store metadata about the cohort definition. Since this table is required it will be created if it it is not supplied.

#### cohort\_attrition:

cohort\_attrition is an optional table that stores attrition information recorded during the cohort generation process such as how many persons were dropped at each step of inclusion rule application. The first column of this table should be cohort\_definition\_id but all other columns currently have no constraints.

#### cohort\_count:

cohort\_count is a option attribute table that records the number of records and the number of unique persons in each cohort in a cohort\_table. It is derived metadata that can be re-derived as long as cohort\_set, the complete list of cohorts in the set, is available. Column names of cohort\_count are: cohort\_definition\_id, number\_records, number\_subjects. This table is required for cohort\_table objects and will be created if not supplied.

#### Value

A cohort\_table object that is a tbl\_sql reference to a cohort table in the write\_schema of an OMOP CDM

#### **Examples**

```
## Not run:
# This function is for developers who are creating cohort_table
# objects in their packages. The function should accept a cdm_reference
# object as the first argument and return a cdm_reference object with the
# cohort table added. The second argument should be `name` which will be
# the prefix for the database tables, the name of the cohort table in the
# database and the name of the cohort table in the cdm object.
# Other optional arguments can be added after the first two.

generateCustomCohort <- function(cdm, name, ...) {

# accept a cdm_reference object as input
checkmate::assertClass(cdm, "cdm_reference")
con <- attr(cdm, "dbcon")

# Create the tables in the database however you like
# All the tables should be prefixed with `name`</pre>
```

read\_cohort\_set 41

```
# The cohort table should be called `name` in the database
  # Create the dplyr table references
  cohort_ref <- dplyr::tbl(con, name)</pre>
  cohort_set <- dplyr::tbl(con, paste0(name, "_set"))</pre>
  cohort_attrition_ref <- dplyr::tbl(con, paste0(name, "_attrition"))</pre>
  cohort_count_ref <- dplyr::tbl(con, paste0(name, "_count"))</pre>
   # add to the cdm
  cdm[[name]] <- cohort_ref</pre>
  # create the generated cohort set object using the constructor
  cdm[[name]] <- new_generated_cohort_set(</pre>
      cdm[[name]],
      cohort_set_ref = cohort_set_ref,
      cohort_attrition_ref = cohort_attrition_ref,
      cohort_count_ref = cohort_count_ref)
   return(cdm)
## End(Not run)
```

read\_cohort\_set

Read a set of cohort definitions into R

## Description

A "cohort set" is a collection of cohort definitions. In R this is stored in a dataframe with cohort\_definition\_id, cohort\_name, and cohort columns. On disk this is stored as a folder with a CohortsToCreate.csv file and one or more json files. If the CohortsToCreate.csv file is missing then all of the json files in the folder will be used, cohort\_definition\_id will be automatically assigned in alphabetical order, and cohort\_name will match the file names.

#### Usage

```
read_cohort_set(path)
readCohortSet(path)
```

#### **Arguments**

path

The path to a folder containing Circe cohort definition json files and optionally a csv file named CohortsToCreate.csv with columns cohortId, cohortName, and jsonPath.

42 recordCohortAttrition

recordCohortAttrition Add attrition reason to a cohort\_table object

#### **Description**

Update the cohort attrition table with new counts and a reason for attrition.

## Usage

```
recordCohortAttrition(cohort, reason, cohortId = NULL)
record_cohort_attrition(cohort, reason, cohortId = NULL)
```

## **Arguments**

cohort A generated cohort set

reason The reason for attrition as a character string

cohortId Cohort definition id of the cohort you want to update the attrition

#### Value

The cohort object with the attributes created or updated.

#### [Experimental]

# **Examples**

```
## Not run:
library(CDMConnector)
library(dplyr)
con <- DBI::dbConnect(duckdb::duckdb(), eunomia_dir())</pre>
cdm <- cdm_from_con(con = con, cdm_schema = "main", write_schema = "main")</pre>
cdm <- generateConceptCohortSet(</pre>
  cdm = cdm, conceptSet = list(pharyngitis = 4112343), name = "new_cohort"
settings(cdm$new_cohort)
cohortCount(cdm$new_cohort)
cohortAttrition(cdm$new_cohort)
cdm$new_cohort <- cdm$new_cohort %>%
  filter(cohort_start_date >= as.Date("2010-01-01"))
cdm$new_cohort <- updateCohortAttributes(</pre>
  cohort = cdm$new_cohort, reason = "Only events after 2010"
)
settings(cdm$new_cohort)
```

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```
cohortCount(cdm$new_cohort)
cohortAttrition(cdm$new_cohort)
## End(Not run)
```

snapshot

Extract CDM metadata

# Description

Extract the name, version, and selected record counts from a cdm.

## Usage

```
snapshot(cdm)
```

## **Arguments**

cdm

A cdm object

#### Value

A named list of attributes about the cdm including selected fields from the cdm\_source table and record counts from the person and observation\_period tables

## **Examples**

```
## Not run:
library(CDMConnector)
con <- DBI::dbConnect(duckdb::duckdb(), eunomia_dir())
cdm <- cdm_from_con(con, "main")
snapshot(cdm)

DBI::dbDisconnect(con, shutdown = TRUE)
## End(Not run)</pre>
```

stow

Collect a list of lazy queries and save the results as files

## **Description**

Collect a list of lazy queries and save the results as files

## Usage

```
stow(cdm, path, format = "parquet")
```

44 summarise\_quantile

#### **Arguments**

cdm A cdm object

path A folder to save the cdm object to

format The file format to use: "parquet" (default), "csv", "feather" or "duckdb".

#### Value

Invisibly returns the cdm input

## **Examples**

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomia_dir())
vocab <- cdm_from_con(con, "main") %>%
    cdm_select_tbl("concept", "concept_ancestor")
stow(vocab, here::here("vocab_tables"))
DBI::dbDisconnect(con, shutdown = TRUE)
## End(Not run)
```

summarise\_quantile

Quantile calculation using dbplyr

## Description

This function provides DBMS independent syntax for quantiles estimation. Can be used by itself or in combination with mutate() when calculating other aggregate metrics (min, max, mean). summarise\_quantile(), summarize\_quantile(), summarizeQuantile() are synonyms.

#### Usage

```
summarise_quantile(.data, x = NULL, probs, name_suffix = "value")
summarize_quantile(.data, x = NULL, probs, name_suffix = "value")
summariseQuantile(.data, x = NULL, probs, nameSuffix = "value")
summarizeQuantile(.data, x = NULL, probs, nameSuffix = "value")
```

## Arguments

```
. data lazy data frame backed by a database query.

x column name whose sample quantiles are wanted.

probs numeric vector of probabilities with values in [0,1].

name_suffix, nameSuffix
```

character; is appended to numerical quantile value as a column name part.

tbl\_group 45

#### **Details**

Implemented quantiles estimation algorithm returns values analogous to quantile{stats} with argument type = 1. See discussion in Hyndman and Fan (1996). Results differ from PERCENTILE\_CONT natively implemented in various DBMS, where returned values are equal to quantile{stats} with default argument type = 7

#### Value

An object of the same type as '.data'

## **Examples**

tbl\_group

CDM table selection helper

## **Description**

The OMOP CDM tables are grouped together and the tbl\_group function allows users to easily create a CDM reference including one or more table groups.

# Usage

```
tbl_group(group)
tblGroup(group)
```

# Arguments

group

A character vector of CDM table groups: "vocab", "clinical", "all", "default", "derived".

46 union\_cohorts

#### **Details**

```
" alt" alt
```

The "default" table group is meant to capture the most commonly used set of CDM tables. Currently the "default" group is: person, observation\_period, visit\_occurrence, visit\_detail, condition\_occurrence, drug\_exposure, procedure\_occurrence, device\_exposure, measurement, observation, death, note, note\_nlp, specimen, fact\_relationship, location, care\_site, provider, payer\_plan\_period, cost, drug\_era, dose\_era, condition\_era, concept, vocabulary, concept\_relationship, concept\_ancestor, concept\_synonym, drug\_strength

#### Value

A character vector of CDM tables names in the groups

## **Examples**

union\_cohorts

Union all cohorts in a single cohort table

## Description

Union all cohorts in a single cohort table

## Usage

```
union_cohorts(x, cohort_definition_id = 1L)
unionCohorts(x, cohort_definition_id = 1L)
```

## **Arguments**

uniqueTableName 47

## Value

A lazy query that when executed will resolve to a new cohort table with one cohort\_definition\_id resulting from the union of all cohorts in the original cohort table

uniqueTableName

Create a unique table name for temp tables

## **Description**

Create a unique table name for temp tables

# Usage

```
uniqueTableName()
unique_table_name()
```

#### Value

A string that can be used as a dbplyr temp table name

validate\_cdm

Validation report for a CDM

## **Description**

Print a short validation report for a cdm object. The validation includes checking that column names are correct and that no tables are empty. A short report is printed to the console. This function is meant for interactive use.

# Usage

```
validate_cdm(cdm)
validateCdm(cdm)
```

# **Arguments**

cdm

A cdm reference object.

#### Value

Invisibly returns the cdm input

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## **Examples**

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb(), eunomia_dir())
cdm <- cdm_from_con(con, cdm_schema = "main")
validate_cdm(cdm)
DBI::dbDisconnect(con)
## End(Not run)</pre>
```

version

Get the CDM version

# Description

Extract the CDM version attribute from a cdm\_reference object

# Usage

```
version(cdm)
```

## **Arguments**

 $\mathsf{cdm}$ 

A cdm object

# Value

```
"5.3" or "5.4"
```

# **Examples**

```
## Not run:
library(CDMConnector)
con <- DBI::dbConnect(duckdb::duckdb(), eunomia_dir())
cdm <- cdm_from_con(con, cdm_schema = "main", write_schema = "main")
version(cdm)

DBI::dbDisconnect(con, shutdown = TRUE)
## End(Not run)</pre>
```

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