

Package ‘CDMConnector’

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Title Connect to an OMOP Common Data Model

Version 2.1.1

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.

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

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

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appendPermanent	<i>Run a dplyr query and add the result set to an existing</i>
-----------------	--

Description

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

Usage

```
appendPermanent(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

Examples

```
## Not run:  
library(CDMConnector)  
  
con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomiaDir())  
concept <- dplyr::tbl(con, "concept")  
  
# 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	<i>as.Date dbplyr translation wrapper</i>
--------	---

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)
```

Arguments

x	an R expression
---	-----------------

Examples

```
## Not run:
con <- DBI::dbConnect(odbc::odbc(), "Oracle")
date_tbl <- dplyr::copy_to(con,
                           data.frame(y = 2000L, m = 10L, d = 10L),
                           name = "tmp",
                           temporary = TRUE)

df <- date_tbl %>%
  dplyr::mutate(date_from_parts = !!asDate(paste0(
    .data$y, "/",
    .data$m, "/",
    .data$d
  ))) %>%
  dplyr::collect()

## End(Not run)
```

benchmarkCDMConnector	<i>Run benchmark of tasks using CDMConnector</i>
-----------------------	--

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(), eunomiaDir())  
cdm <- cdmFromCon(con, cdmSchema = "main", writeSchema = "main")  
benchmarkCDMConnector(cdm)  
  
DBI::dbDisconnect(con, shutdown = TRUE)  
  
## End(Not run)
```

cdmCon

Get underlying database connection

Description

Get underlying database connection

Usage

cdmCon(cdm)

Arguments

cdm A cdm reference object created by cdmFromCon

Value

A reference to the database containing tables in the cdm reference

Examples

```
## Not run:  
con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomiaDir())  
  
cdm <- cdmFromCon(con = con, cdmName = "Eunomia",  
                    cdmSchema = "main", writeSchema = "main")  
  
cdmCon(cdm)  
  
DBI::dbDisconnect(con)
```

```
## End(Not run)
```

`cdmDisconnect.db_cdm` *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

```
## S3 method for class 'db_cdm'
cdmDisconnect(cdm, dropWriteSchema = FALSE, ...)
```

Arguments

<code>cdm</code>	cdm reference
<code>dropWriteSchema</code>	Whether to drop tables in the writeSchema
<code>...</code>	Not used

`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.

Usage

```
cdmFlatten(
  cdm,
  domain = c("condition_occurrence", "drug_exposure", "procedure_occurrence"),
  includeConceptName = TRUE
)
```

Arguments

cdm	A cdm_reference object
domain	Domains to include. Must be a subset of "condition_occurrence", "drug_exposure", "procedure_occurrence", "measurement", "visit_occurrence", "death", "observation"
includeConceptName	Should concept_name and type_concept_name be include in the output table? TRUE (default) or FALSE

Value

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

Examples

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

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

cdm <- cdmFromCon(con, cdmSchema = "main")

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>
#> 1       2          40213201 1986-09-09 1986-09-09  5.81e5 drug   pneumo <NA>
#> 2       18         4116491  1997-11-09 1998-01-09  3.20e4 cond   Escher <NA>
#> 3       18         40213227 2017-01-04 2017-01-04  5.81e5 drug   tetanu <NA>
#> 4       42         4156265  1974-06-13 1974-06-27  3.20e4 cond   Facial <NA>
#> 5       18         40213160 1966-02-23 1966-02-23  5.81e5 drug   poliov <NA>
#> 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 cond   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      2          4163872  1993-03-29 1993-03-29  3.80e7 proce  Plain  <NA>
#> # ... 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)
```

cdmFromCon*Create a CDM reference object from a database connection*

Description

Create a CDM reference object from a database connection

Usage

```
cdmFromCon(
  con,
  cdmSchema,
  writeSchema,
  cohortTables = NULL,
  cdmVersion = NULL,
  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.
cdmSchema	The schema where the OMOP CDM tables are located. Defaults to NULL.
writeSchema	An optional schema in the CDM database that the user has write access to.
cohortTables	A character vector listing the cohort table names to be included in the CDM object.
cdmVersion	The version of the OMOP CDM. Can be "5.3", "5.4", or NULL (default). If NULL we will attempt to automatically determine the cdm version using the cdm_source table and heuristics.
cdmName	The name of the CDM. If NULL (default) the cdm_source_name . field in the CDM_SOURCE table will be used.
achillesSchema	An optional schema in the CDM database that contains achilles tables.
.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.
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

cdmFromCon creates a new cdm reference object from a DBI 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.

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 writePrefix. This is a short character string that will be added to any tables created in the writeSchema effectively a namespace in the schema just for your analysis. If the write_schema is a shared between multiple users setting a unique write_prefix ensures you do not overwrite existing tables and allows you to easily clean up tables by dropping all tables that start with the prefix.

Value

A list of dplyr database table references pointing to CDM tables

Examples

```
## Not run:  
library(CDMConnector)  
con <- DBI::dbConnect(duckdb::duckdb(), eunomiaDir())  
  
# minimal example  
cdm <- cdmFromCon(con,  
                     cdmSchema = "main",  
                     writeSchema = "scratch")  
  
# write prefix is optional but recommended if write_schema is shared  
cdm <- cdmFromCon(con,  
                     cdmSchema = "main",  
                     writeSchema = "scratch",  
                     writePrefix = "tmp_")  
  
# Some database systems use catalogs or compound schemas.  
# These can be specified as follows:  
cdm <- cdmFromCon(con,  
                     cdmSchema = "catalog.main",  
                     writeSchema = "catalog.scratch",  
                     writePrefix = "tmp_")  
  
cdm <- cdmFromCon(con,  
                     cdmSchema = c("my_catalog", "main"),  
                     writeSchema = c("my_catalog", "scratch"),  
                     writePrefix = "tmp_")  
  
cdm <- cdmFromCon(con,  
                     cdmSchema = c(catalog = "my_catalog", schema = "main"),  
                     writeSchema = c(catalog = "my_catalog", schema = "scratch"))
```

```

        writePrefix = "tmp_")

DBI::dbDisconnect(con)

## End(Not run)

```

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")
```

Arguments

<code>cdm</code>	A <code>cdm_reference</code> object.
<code>n</code>	Number of persons to include in the <code>cdm</code> .
<code>seed</code>	Seed for the random number generator.
<code>name</code>	Name of the table that will contain the sample of persons.

Value

A modified `cdm_reference` object where all clinical tables are lazy queries pointing to subset

Examples

```

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

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

cdm <- cdmFromCon(con, cdmSchema = "main")

cdmSampled <- cdmSample(cdm, n = 2)

cdmSampled$person %>%
  select(person_id)
#> # Source:   SQL [2 x 1]
#> # Database: DuckDB 0.6.1

```

```
#>   person_id
#>     <dbl>
#> 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)
```

Arguments

cdm	A <code>cdm_reference</code> object
personId	A numeric vector of person IDs to include in the <code>cdm</code>

Value

A modified `cdm_reference` object where all clinical tables are lazy queries pointing to subset

Examples

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

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

cdm <- cdmFromCon(con, cdmSchema = "main")

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

cdm2$person %>%
  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
#> 3      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)
```

Arguments

<code>cdm</code>	A <code>cdm_reference</code> object
<code>cohortTable</code>	The name of a cohort table in the <code>cdm</code> reference
<code>cohortId</code>	IDs of the cohorts that we want to subset from the cohort table. If <code>NULL</code> (default) all cohorts in cohort table are considered.
<code>verbose</code>	Should subset messages be printed? <code>TRUE</code> or <code>FALSE</code> (default)

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(), eunomiaDir())

cdm <- cdmFromCon(con, cdmSchema = "main", writeSchema = "main")

# generate a cohort
path <- system.file("cohorts2", mustWork = TRUE, package = "CDMConnector")

cohortSet <- readCohortSet(path) %>%
```

```
filter(cohort_name == "GIBleed_male")

# subset cdm to persons in the generated cohort
cdm <- generateCohortSet(cdm, cohortSet = cohortSet, name = "gibleed")

cdmGiBleed <- cdmSubsetCohort(cdm, cohortTable = "gibleed")

cdmGiBleed$person %>%
  tally()
#> # Source:   SQL [1 x 1]
#> # Database: DuckDB 0.6.1
#>      n
#>    <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

Usage

```
cdmWriteSchema(cdm)
```

Arguments

cdm	A cdm reference object created by cdmFromCon
-----	--

Value

The database write schema

Examples

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomiaDir())

cdm <- cdmFromCon(con = con, cdmName = "Eunomia",
                     cdmSchema = "main", writeSchema = "main")

cdmWriteSchema(cdm)

DBI::dbDisconnect(con)

## End(Not run)
```

`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.

Usage

```
computeQuery(
  x,
  name = uniqueTableName(),
  temporary = TRUE,
  schema = NULL,
  overwrite = TRUE,
  ...
)
```

Arguments

<code>x</code>	A <code>dplyr</code> query
<code>name</code>	The name of the table to create.
<code>temporary</code>	Should the table be temporary: TRUE (default) or FALSE
<code>schema</code>	The schema where the table should be created. Ignored if <code>temporary = TRUE</code> .
<code>overwrite</code>	Should the table be overwritten if it already exists: TRUE (default) or FALSE Ignored if <code>temporary = TRUE</code> .
<code>...</code>	Further arguments passed on the <code>dplyr::compute</code>

Value

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

Examples

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

con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomiaDir())
cdm <- cdmFromCon(con, "main")

# 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 %>%
  dplyr::count(domain_id == "Drug") %>%
  computeQuery("tmp_table", temporary = FALSE, schema = "main")

DBI::dbDisconnect(con, shutdown = TRUE)

## End(Not run)
```

copyCdmTo

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. `copyCdmTo` 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

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

Arguments

<code>con</code>	A DBI database connection created by <code>DBI::dbConnect</code>
<code>cdm</code>	A <code>cdm</code> reference object created by <code>CDMConnector::cdmFromCon</code> or <code>CDMConnector::cdm_from_con</code>
<code>schema</code>	schema name in the remote database where the user has write permission
<code>overwrite</code>	Should the cohort table be overwritten if it already exists? TRUE or FALSE (default)

Value

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

dateadd	<i>Add days or years to a date in a dplyr query</i>
---------	---

Description

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

Usage

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

Arguments

date	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

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb())
date_tbl <- dplyr::copy_to(con, data.frame(date1 = as.Date("1999-01-01")),
                           name = "tmpdate", overwrite = TRUE, temporary = TRUE)

df <- date_tbl %>%
  dplyr::mutate(date2 = !!dateadd("date1", 1, interval = "year")) %>%
  dplyr::collect()

DBI::dbDisconnect(con, shutdown = TRUE)

## End(Not run)
```

datediff	<i>Compute the difference between two days</i>
----------	--

Description

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

Usage

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

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

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb())
date_tbl <- dplyr::copy_to(con, data.frame(date1 = as.Date("1999-01-01")),
                           name = "tmpdate", overwrite = TRUE, temporary = TRUE)

df <- date_tbl %>%
  dplyr::mutate(date2 = !!dateadd("date1", 1, interval = "year")) %>%
  dplyr::mutate(dif_years = !!datediff("date1", "date2", interval = "year")) %>%
  dplyr::collect()

DBI::dbDisconnect(con, shutdown = TRUE)

## End(Not run)
```

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.

Examples

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb(), ":memory:")
date_tbl <- dplyr::copy_to(con,
                           data.frame(birth_date = as.Date("1993-04-19")),
                           name = "tmp",
                           temporary = TRUE)
df <- date_tbl %>%
  dplyr::mutate(year = !!datepart("birth_date", "year")) %>%
  dplyr::mutate(month = !!datepart("birth_date", "month")) %>%
  dplyr::mutate(day = !!datepart("birth_date", "day")) %>%
  dplyr::collect()
DBI::dbDisconnect(con, shutdown = TRUE)

## End(Not run)
```

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

Examples

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomiaDir())
cdm <- cdmFromCon(con)
dbms(cdm)
dbms(con)

## End(Not run)
```

dbSource	<i>Create a source for a cdm in a database.</i>
----------	---

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	<i>Download Eunomia data files</i>
---------------------	------------------------------------

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  
)
```

Arguments

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

Value

Invisibly returns the destination if the download was successful.

Examples

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

## End(Not run)
```

dropTable.db_cdm

Drop table from a database backed cdm object

Description

Tables will be dropped from the write schema of the cdm.

Usage

```
## S3 method for class 'db_cdm'
dropTable(cdm, name)
```

Arguments

cdm	a cdm_reference object
name	A character vector of table names to be dropped

eunomiaDir

Create a copy of an example OMOP CDM dataset

Description

Eunomia is an OHDSI project that provides several example OMOP CDM datasets for testing and development. This function creates a copy of a Eunomia database in **duckdb** 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")
)
```

Arguments

datasetName	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-sn1-10k", "synthea-surgery-10k", "synthea-total_joint_replacement-10k", "synthea-veteran_prostate_cancer-10k", "synthea-veterans-10k", "synthea-weight_loss-10k", "empty_cdm", "synpuf-1k"
cdmVersion	The OMOP CDM version. Must be "5.3" or "5.4".
databaseFile	The full path to the new copy of the example CDM dataset.

Details

Most of the Eunomia datasets available in CDMConnector are from the Synthea project. Synthea is an open-source synthetic patient generator that models the medical history of synthetic patients. The Synthea datasets are generated using the Synthea tool and then converted to the OMOP CDM format using the OHDSI ETL-Synthea project <https://ohdsi.github.io/ETL-Synthea/>. Currently the synthea datasets are only available in the OMOP CDM v5.3 format. See <https://synthetichealth.github.io/synthea/> for details on the Synthea project.

In addition to Synthea, the Eunomia project provides the CMS Synthetic Public Use Files (SynPUFs) in both 5.3 and 5.4 OMOP CDM formats. This data is synthetic US Medicare claims data mapped to OMOP CDM format. The OMOP CDM has a set of optional metadata tables, called Achilles tables, that include pre-computed analytics about the entire dataset such as record and person counts. The Eunomia Synpuf datasets include the Achilles tables.

Eunomia also provides empty cdms that can be used as a starting point for creating a new example CDM. This is useful for creating test data for studies or analytic packages. The empty CDM includes the vocabulary tables and all OMOP CDM tables but the clinical tables are empty and need to be populated with data. For additional information on creating small test CDM datasets see <https://ohdsi.github.io/omock/> and <https://darwin-eu.github.io/TestGenerator/>.

To contribute synthetic observational health data to the Eunomia project please open an issue at <https://github.com/OHDSI/Eunomia/issues/>

Setup: To use the eunomiaDir function please set the EUNOMIA_DATA_FOLDER in your .Renviron file to a folder on your computer where the datasets will be downloaded to. This file can be opened by calling `usethis::edit_r_environ()`.

Value

The file path to the new Eunomia dataset copy

Examples

```
## Not run:

# The defaults GiBleed dataset is a small dataset that is useful for testing
library(CDMConnector)
con <- DBI::dbConnect(duckdb::duckdb(), eunomiaDir())
```

```

cdm <- cdmFromCon(con, "main", "main")
cdmDisconnect(cdm)

# Synpuf datasets include the Achilles tables
con <- DBI::dbConnect(duckdb::duckdb(), eunomiaDir("synpuf-1k", "5.3"))
cdm <- cdmFromCon(con, "main", "main", achillesSchema = "main")
cdmDisconnect(cdm)

# Currently the only 5.4 dataset is synpuf-1k
con <- DBI::dbConnect(duckdb::duckdb(), eunomiaDir("synpuf-1k", "5.4"))
cdm <- cdmFromCon(con, "main", "main", achillesSchema = "main")
cdmDisconnect(cdm)

## End(Not run)

```

eunomiaIsAvailable *Has the Eunomia dataset been cached?*

Description

Has the Eunomia dataset been cached?

Usage

```
eunomiaIsAvailable(datasetName = "GiBleed", cdmVersion = "5.3")
```

Arguments

datasetName	Name of the Eunomia dataset to check. Defaults to "GiBleed".
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 FALSE otherwise

exampleDatasets *List the available example CDM datasets*

Description

List the available example CDM datasets

Usage

```
exampleDatasets()
```

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 <- cdmFromCon(con)

## End(Not run)
```

generateCohortSet

Generate a cohort set on a cdm object

Description

A "cohort_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

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 its attributes may no longer be accurate.

Usage

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

Arguments

cdm	A cdm reference created by CDMConnector. write_schema must be specified.
cohortSet	A cohortSet dataframe created with readCohortSet()
name	Name of the cohort table to be created. This will also be used as a prefix for the cohort attribute tables. This must be a lowercase character string that starts with a letter and only contains letters, numbers, and underscores.
computeAttrition	Should attrition be computed? TRUE (default) or FALSE
overwrite	Should the cohort table be overwritten if it already exists? TRUE (default) or FALSE

Examples

```
## Not run:
library(CDMConnector)
con <- DBI::dbConnect(duckdb::duckdb(), eunomiaDir())
cdm <- cdmFromCon(con,
                    cdmSchema = "main",
                    writeSchema = "main")

cohortSet <- readCohortSet(system.file("cohorts2", package = "CDMConnector"))
cdm <- generateCohortSet(cdm, cohortSet, name = "cohort")

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
)
```

Arguments

<code>cdm</code>	A cdm reference object created by <code>CDMConnector::cdmFromCon</code> or <code>CDMConnector::cdm_from_con</code>
<code>conceptSet</code>	A named list of numeric vectors or a Concept Set Expression created <code>omopgenerics::newConceptSetExp</code>
<code>name</code>	The name of the new generated cohort table as a character string
<code>limit</code>	Include "first" (default) or "all" occurrences of events in the cohort <ul style="list-style-type: none"> "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.
<code>requiredObservation</code>	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.
<code>end</code>	How should the <code>cohort_end_date</code> be defined? <ul style="list-style-type: none"> "<code>observation_period_end_date</code>" (default): The earliest <code>observation_period_end_date</code> after the event start date numeric scalar: A fixed number of days from the event start date "<code>event_end_date</code>": The event end date. If the event end date is not populated then the event start date will be used
<code>subsetCohort</code>	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.
<code>subsetCohortId</code>	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.
<code>overwrite</code>	Should the cohort table be overwritten if it already exists? <code>TRUE</code> (default) or <code>FALSE</code> .

Value

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

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)
```

Arguments

<code>schema</code>	A schema name as a character string
<code>table</code>	A table name as character string
<code>dbms</code>	The name of the database management system as returned by dbms(connection)

Value

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

listTables*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

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

Arguments

<code>con</code>	A DBI connection to a database
<code>schema</code>	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 = eunomiaDir())
listTables(con, schema = "main")

## End(Not run)
```

readCohortSet

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

```
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.
------	---

requireEunomia

Require eunomia to be available. The function makes sure that you can later create a eunomia database with eunomiaDir().

Description

Require eunomia to be available. The function makes sure that you can later create a eunomia database with eunomiaDir().

Usage

```
requireEunomia(datasetName = "GiBleed", cdmVersion = "5.3")
```

Arguments

datasetName	Name of the Eunomia dataset to check. Defaults to "GiBleed".
cdmVersion	Version of the Eunomia dataset to check. Must be "5.3" or "5.4".

Value

Path to eunomia database.

<code>snapshot</code>	<i>Extract CDM metadata</i>
-----------------------	-----------------------------

Description

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

Usage

```
snapshot(cdm)
```

Arguments

<code>cdm</code>	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(), eunomiaDir())
cdm <- cdmFromCon(con, "main")
snapshot(cdm)

DBI::dbDisconnect(con, shutdown = TRUE)

## End(Not run)
```

<code>summariseQuantile</code>	<i>Quantile calculation using dbplyr</i>
--------------------------------	--

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()`, `summariseQuantile()` and `summarizeQuantile()` are synonyms.

Usage

```
summariseQuantile(.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].
nameSuffix	character; is appended to numerical quantile value as a column name part.

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

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb())
mtcars_tbl <- dplyr::copy_to(con, mtcars, name = "tmp", overwrite = TRUE, temporary = TRUE)

df <- mtcars_tbl %>%
  dplyr::group_by(cyl) %>%
  dplyr::mutate(mean = mean(mpg, na.rm = TRUE)) %>%
  summariseQuantile(mpg, probs = c(0, 0.2, 0.4, 0.6, 0.8, 1),
                     nameSuffix = "quant") %>%
  dplyr::collect()

DBI::dbDisconnect(con, shutdown = TRUE)

## End(Not run)
```

summariseQuantile2 *Quantile calculation using dbplyr*

Description

This function provides DBMS independent syntax for quantile estimation. Some database systems do not have a quantile function. The SQL generated by `summarizeQuantile2` should work on all supported database systems. This function can be added to a `dplyr` pipeline and adds an additional query to the input. No computation is triggered by `summarizeQuantile2` if the input is a `tbl` reference to a database table.

Usage

```
summariseQuantile2(.data, x, probs, nameSuffix = "{x}")
```

Arguments

.data	lazy data frame backed by a database query created by <code>dplyr::tbl()</code> .
x	A string vector of column names whose sample quantiles are wanted.
probs	A numeric vector of probabilities with values in [0,1].
nameSuffix	A single character character string, evaluated by <code>glue::glue()</code> that is appended to numerical quantile value as a column name part.

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

[Experimental]

Value

A lazy query with quantile calculation added. The result (after computation) will have one row per combination of grouping variables and one column for every variable/quantile combination. (see examples)

Examples

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb())
mtcars_tbl <- dplyr::copy_to(con, mtcars, name = "tmp", overwrite = TRUE, temporary = TRUE)

# quantiles for a single column
mtcars_tbl %>%
  dplyr::group_by(cyl) %>%
  dplyr::mutate(mean = mean(mpg, na.rm = TRUE)) %>%
  summariseQuantile2("mpg", probs = c(0, 0.2, 0.4, 0.6, 0.8, 1), nameSuffix = "quant") %>%
  dplyr::collect()

#> cyl  p0_quant p20_quant p40_quant p60_quant p80_quant p100_quant
#>   6    17.8     18.1     19.2      21       21     21.4
#>   8    10.4     13.3     15       15.5     17.3    19.2
#>   4    21.4     22.8     24.4     27.3     30.4    33.9

# multiple columns
mtcars_tbl %>%
  dplyr::group_by(cyl) %>%
  dplyr::mutate(mean = mean(mpg, na.rm = TRUE)) %>%
  summariseQuantile2(c("mpg", "hp", "wt"), probs = c(0.2, 0.8), nameSuffix = "{x}_quant") %>%
  dplyr::collect()
```

```

#> cyl p20_mpg_quant p80_mpg_quant p20_hp_quant p80_hp_quant p20_wt_quant p80_wt_quant
#>   4      22.8      30.4      65      97     1.84     2.78
#>   6      18.1       21     110     123     2.77     3.44
#>   8      13.3      17.3     175     245     3.44     5.25

DBI::dbDisconnect(con, shutdown = TRUE)

## End(Not run)

```

tblGroup *CDM table selection helper*

Description

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

Usage

tblGroup(group)

Arguments

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

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

```
cdm <- cdmFromCon(con, cdmName = "test", cdmSchema = "public") %>%
  cdmSelectTbl(tblGroup("vocab"))

## End(Not run)
```

version*Get the CDM version***Description**

Extract the CDM version attribute from a `cdm_reference` object

Usage

```
version(cdm)
```

Arguments

<code>cdm</code>	A <code>cdm</code> object
------------------	---------------------------

Value

"5.3" or "5.4"

Examples

```
## Not run:
library(CDMConnector)
con <- DBI::dbConnect(duckdb::duckdb(), eunomiaDir())
cdm <- cdmFromCon(con, cdmSchema = "main", writeSchema = "main")
version(cdm)

DBI::dbDisconnect(con, shutdown = TRUE)

## End(Not run)
```

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