Package 'omopgenerics'

December 5, 2024

```
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    Description Provides definitions of core classes and methods used by analytic pipelines that query the OMOP (Observational Medical Outcomes Partnership) common data model.
```

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achillesColumns

Required columns for each of the achilles result tables

Description

Required columns for each of the achilles result tables

Usage

```
achillesColumns(table, version = "5.3", onlyRequired = lifecycle::deprecated())
```

achillesTables 5

Arguments

table Table for which to see the required columns. One of "achilles_analysis", "achilles_results",

or "achilles_results_dist".

version Version of the OMOP Common Data Model.

onlyRequired deprecated.

Value

Character vector with the column names

Examples

```
library(omopgenerics)
achillesColumns("achilles_analysis")
achillesColumns("achilles_results")
achillesColumns("achilles_results_dist")
```

achillesTables

Names of the tables that contain the results of achilles analyses

Description

Names of the tables that contain the results of achilles analyses

Usage

```
achillesTables(version = "5.3")
```

Arguments

version

Version of the OMOP Common Data Model.

Value

Names of the tables that are contain the results from the achilles analyses

```
library(omopgenerics)
achillesTables()
```

6 additionalColumns

additionalColumns

Identify variables in additional_name column

Description

Identifies and returns the unique values in additional_name column.

Usage

```
additionalColumns(result)
```

Arguments

result

A tibble.

Value

Unique values of the additional name column.

```
{
 library(dplyr)
 library(omopgenerics)
 x <- tibble(
    "result_id" = as.integer(c(1, 2)),
    "cdm_name" = c("cprd", "eunomia"),
    "group_name" = "cohort_name",
    "group_level" = "my_cohort",
    "strata_name" = "sex",
    "strata_level" = "male",
    "variable_name" = "Age group",
    "variable_level" = "10 to 50",
    "estimate_name" = "count",
    "estimate_type" = "numeric",
    "estimate_value" = "5",
    "additional_name" = "overall",
    "additional_level" = "overall"
 ) |>
   newSummarisedResult(settings = tibble(
      "result_id" = c(1, 2), "custom" = c("A", "B")
 x |> additionalColumns()
```

addSettings 7

 ${\tt addSettings}$

Add settings columns to a <summarised_result> object

Description

Add settings columns to a <summarised_result> object

Usage

```
addSettings(result, settingsColumn = settingsColumns(result))
```

Arguments

result A <summarised_result> object.

settingsColumn Settings to be added as columns, by default settingsColumns(result) will be added. If NULL or empty character vector, no settings will be added.

Value

A <summarised_result> object with the added setting columns.

```
{
 library(dplyr)
 library(omopgenerics)
 x <- tibble(</pre>
    "result_id" = as.integer(c(1, 2)),
    "cdm_name" = c("cprd", "eunomia"),
    "group_name" = "cohort_name",
    "group_level" = "my_cohort",
    "strata_name" = "sex",
    "strata_level" = "male",
    "variable_name" = "Age group",
    "variable_level" = "10 to 50",
    "estimate_name" = "count",
    "estimate_type" = "numeric",
    "estimate_value" = "5",
    "additional_name" = "overall",
    "additional_level" = "overall"
 ) |>
    newSummarisedResult(settings = tibble(
      "result_id" = c(1, 2), "custom" = c("A", "B")
   ))
 Χ
 x |> addSettings()
```

8 assertCharacter

}

assertCharacter

Assert that an object is a character and fulfill certain conditions.

Description

Assert that an object is a character and fulfill certain conditions.

Usage

```
assertCharacter(
    x,
    length = NULL,
    na = FALSE,
    null = FALSE,
    unique = FALSE,
    named = FALSE,
    minNumCharacter = 0,
    call = parent.frame(),
    msg = NULL
)
```

Arguments

x Variable to check.

length Required length. If NULL length is not checked.

na Whether it can contain NA values.

null Whether it can be NULL.

unique Whether it has to contain unique elements.

named Whether it has to be named.

minNumCharacter

Minimum number of characters that all elements must have.

call Call argument that will be passed to cli error message.

msg Custom error message.

assertChoice 9

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Assert that an object is within a certain oprtions.

Description

Assert that an object is within a certain oprtions.

Usage

```
assertChoice(
    x,
    choices,
    length = NULL,
    na = FALSE,
    null = FALSE,
    unique = FALSE,
    named = FALSE,
    call = parent.frame(),
    msg = NULL
)
```

Arguments

X	Variable to check.
choices	Options that x is allowed to be.
length	Required length. If NULL length is not checked.
na	Whether it can contain NA values.
null	Whether it can be NULL.
unique	Whether it has to contain unique elements.
named	Whether it has to be named.
call	Call argument that will be passed to cli error message.
msg	Custom error message.

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Assert that an object has a certain class.

Description

Assert that an object has a certain class.

10 assertDate

Usage

```
assertClass(
    x,
    class,
    length = NULL,
    null = FALSE,
    all = FALSE,
    extra = TRUE,
    call = parent.frame(),
    msg = NULL
)
```

Arguments

X	To check.
class	Expected class or classes.
length	Required length. If NULL length is not checked.
null	Whether it can be NULL.
all	Whether it should have all the classes or only at least one of them.
extra	Whether the object can have extra classes.
call	Call argument that will be passed to cli.
msg	Custom error message.

assertDate Assert Date

Description

Assert Date

Usage

```
assertDate(
    X,
    length = NULL,
    na = FALSE,
    null = FALSE,
    unique = FALSE,
    named = FALSE,
    call = parent.frame(),
    msg = NULL
)
```

assertList 11

Arguments

Х Expression to check. length Required length. na

Whether it can contain NA values.

null Whether it can be NULL.

unique Whether it has to contain unique elements.

named Whether it has to be named.

call Call argument that will be passed to cli error message.

Custom error message. msg

Value

X

assertList

Assert that an object is a list.

Description

Assert that an object is a list.

Usage

```
assertList(
  length = NULL,
 na = FALSE,
  null = FALSE,
  unique = FALSE,
  named = FALSE,
  class = NULL,
  call = parent.frame(),
 msg = NULL
)
```

Arguments

Variable to check. Х

length Required length. If NULL length is not checked.

Whether it can contain NA values. na

null Whether it can be NULL.

Whether it has to contain unique elements. unique

Whether it has to be named. named

12 assertLogical

class Class that the elements must have.

call Call argument that will be passed to cli error message.

msg Custom error message.

assertLogical

Assert that an object is a logical.

Description

Assert that an object is a logical.

Usage

```
assertLogical(
    X,
    length = NULL,
    na = FALSE,
    null = FALSE,
    named = FALSE,
    call = parent.frame(),
    msg = NULL
)
```

Arguments

x Variable to check.

length Required length. If NULL length is not checked.

na Whether it can contain NA values.

null Whether it can be NULL.

named Whether it has to be named.

call Call argument that will be passed to cli error message.

msg Custom error message.

assertNumeric 13

assertNumeric

Assert that an object is a numeric.

Description

Assert that an object is a numeric.

Usage

```
assertNumeric(
    x,
    integerish = FALSE,
    min = -Inf,
    max = Inf,
    length = NULL,
    na = FALSE,
    null = FALSE,
    unique = FALSE,
    named = FALSE,
    call = parent.frame(),
    msg = NULL
)
```

Arguments

Χ

msg

integerish	Whether it has to be an integer
min	Minimum value that the object can be.
max	Maximum value that the object can be.
length	Required length. If NULL length is not checked.
na	Whether it can contain NA values.
null	Whether it can be NULL.
unique	Whether it has to contain unique elements.
named	Whether it has to be named.
call	Call argument that will be passed to cli error message.

Custom error message.

Variable to check.

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assertTable

Assert that an object is a table.

Description

Assert that an object is a table.

Usage

```
assertTable(
    x,
    class = NULL,
    numberColumns = NULL,
    numberRows = NULL,
    columns = character(),
    allowExtraColumns = TRUE,
    null = FALSE,
    unique = FALSE,
    call = parent.frame(),
    msg = NULL
)
```

Arguments

x Variable to check.

class A class that the table must have: "tbl", "data.fram", "tbl_sql", ...

numberColumns Number of columns that it has to contain.

numberRows Number of rows that it has to contain.

columns Name of the columns required.

allowExtraColumns

Whether extra columns are allowed.

null Whether it can be NULL.

unique Whether it has to contain unique rows.

call Call argument that will be passed to cli error message.

msg Custom error message.

assertTrue 15

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Assert that an expression is TRUE.

Description

Assert that an expression is TRUE.

Usage

```
assertTrue(x, null = FALSE, call = parent.frame(), msg = NULL)
```

Arguments

null Whether it can be NULL.

call Call argument that will be passed to cli error message.

msg Custom error message.

attrition

Get attrition from an object.

Description

Get attrition from an object.

Usage

```
attrition(x)
```

Arguments

Х

An object for which to get an attrition summary.

Value

A table with the attrition.

16 attrition.cohort_table

```
attrition.cohort_table
```

Get cohort attrition from a cohort_table object.

Description

Get cohort attrition from a cohort_table object.

Usage

```
## S3 method for class 'cohort_table'
attrition(x)
```

Arguments

Χ

A cohort_table

Value

A table with the attrition.

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
person <- tibble(</pre>
  person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
  race_concept_id = 0, ethnicity_concept_id = 0
)
observation_period <- tibble(</pre>
  observation_period_id = 1, person_id = 1,
  observation_period_start_date = as.Date("2000-01-01"),
  observation_period_end_date = as.Date("2023-12-31"),
  period_type_concept_id = 0
cohort <- tibble(</pre>
  cohort_definition_id = c(1, 1, 1, 2),
  subject_id = 1,
 {\tt cohort\_start\_date = as.Date(c("2020-01-01", "2021-01-01", "2022-01-01", "2022-01-01")),}
  {\tt cohort\_end\_date = as.Date(c("2020-01-01", "2021-01-01", "2022-01-01", "2022-01-01")),}
cdm <- cdmFromTables(</pre>
  tables = list("person" = person, "observation_period" = observation_period),
  cdmName = "my_example_cdm",
  cohortTables = list("cohort1" = cohort)
)
attrition(cdm$cohort1)
```

bind 17

bind

Bind two or more objects of the same class.

Description

Bind two or more objects of the same class.

Usage

```
bind(...)
```

Arguments

... Objects to bind.

Value

New object.

 $\verb|bind.cohort_table|$

Bind two or more cohort tables

Description

Bind two or more cohort tables

Usage

```
## S3 method for class 'cohort_table'
bind(..., name)
```

Arguments

... Generated cohort set objects to bind. At least two must be provided.

name Name of the new generated cohort set.

Value

The cdm object with a new generated cohort set containing all of the cohorts passed.

Examples

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
cohort1 <- tibble(</pre>
  cohort_definition_id = 1,
  subject_id = 1:3,
  cohort_start_date = as.Date("2010-01-01"),
  cohort_end_date = as.Date("2010-01-05")
cohort2 <- tibble(</pre>
  cohort_definition_id = c(2, 2, 3, 3, 3),
  subject_id = c(1, 2, 3, 1, 2),
  cohort_start_date = as.Date("2010-01-01"),
  cohort_end_date = as.Date("2010-01-05")
)
cdm <- cdmFromTables(</pre>
  tables = list(
    "person" = tibble(
      person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
      race_concept_id = 0, ethnicity_concept_id = 0
    ),
    "observation_period" = tibble(
      observation_period_id = 1:3, person_id = 1:3,
      observation_period_start_date = as.Date("2000-01-01"),
      observation_period_end_date = as.Date("2023-12-31"),
      period_type_concept_id = 0
    )
  ),
  cdmName = "mock",
  cohortTables = list("cohort1" = cohort1, "cohort2" = cohort2)
)
cdm <- bind(cdm$cohort1, cdm$cohort2, name = "cohort3")</pre>
settings(cdm$cohort3)
cdm$cohort3
```

bind.summarised_result

Bind two or summarised_result objects

Description

Bind two or summarised_result objects

Usage

```
## S3 method for class 'summarised_result'
bind(...)
```

cdmFromTables 19

Arguments

... summarised_result objects

Value

A summarised_result object the merged objects.

Examples

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
cdm <- cdmFromTables(</pre>
  tables = list(
    "person" = tibble(
      person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
      race_concept_id = 0, ethnicity_concept_id = 0
    "observation_period" = tibble(
      observation_period_id = 1:3, person_id = 1:3,
      observation_period_start_date = as.Date("2000-01-01"),
      observation_period_end_date = as.Date("2023-12-31"),
      period_type_concept_id = 0
    )
  ),
  cdmName = "mock",
  cohortTables = list("cohort1" = tibble(
    cohort_definition_id = 1,
    subject_id = 1:3,
    cohort_start_date = as.Date("2010-01-01"),
    cohort_end_date = as.Date("2010-01-05")
  ))
)
result1 <- summary(cdm)</pre>
result2 <- summary(cdm$cohort1)</pre>
mergedResult <- bind(result1, result2)</pre>
mergedResult
```

 ${\tt cdmFromTables}$

Create a cdm object from local tables

Description

Create a cdm object from local tables

20 cdmName

Usage

```
cdmFromTables(tables, cdmName, cohortTables = list(), cdmVersion = NULL)
```

Arguments

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

cdmName Name of the cdm object.

cohortTables List of tables that contains cohort, cohort_set and cohort_attrition can be pro-

vided as attributes.

cdmVersion Version of the cdm_reference

Value

A cdm_reference object.

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("2023-12-31"),
    period_type_concept_id = 0
)
cdm <- cdmFromTables(
    tables = list("person" = person, "observation_period" = observation_period),
    cdmName = "test"
)</pre>
```

cdmName

Get the name of a cdm_reference associated object

Description

Get the name of a cdm_reference associated object

Usage

```
cdmName(x)
```

cdmReference 21

Arguments

Х

A cdm_reference or cdm_table object.

Value

Name of the cdm_reference.

Examples

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
cdm <- cdmFromTables(</pre>
  tables = list(
    "person" = tibble(
      person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
      race_concept_id = 0, ethnicity_concept_id = 0
    "observation_period" = tibble(
      observation_period_id = 1:3, person_id = 1:3,
      observation_period_start_date = as.Date("2000-01-01"),
      observation_period_end_date = as.Date("2023-12-31"),
      period_type_concept_id = 0
  ),
  cdmName = "mock"
)
cdmName(cdm)
cdmName(cdm$person)
```

 ${\it cdmReference}$

Get the cdm_reference *of a* cdm_table.

Description

Get the cdm_reference of a cdm_table.

Usage

```
cdmReference(table)
```

Arguments

table

A cdm_table.

22 cdmSelect

Value

A cdm_reference.

Examples

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
cdm <- cdmFromTables(</pre>
  tables = list(
    "person" = tibble(
      person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
      race_concept_id = 0, ethnicity_concept_id = 0
    ),
    "observation_period" = tibble(
      observation_period_id = 1:3, person_id = 1:3,
      observation_period_start_date = as.Date("2000-01-01"),
      observation_period_end_date = as.Date("2023-12-31"),
      period_type_concept_id = 0
  ),
  cdmName = "mock"
cdmReference(cdm$person)
```

cdmSelect

Restrict the cdm object to a subset of tables.

Description

Restrict the cdm object to a subset of tables.

Usage

```
cdmSelect(cdm, ...)
```

Arguments

cdm A cdm_reference object.

... Selection of tables to use, it supports tidyselect expressions.

Value

A cdm_reference with only the specified tables.

cdmSource 23

Examples

```
cdm <- emptyCdmReference("my cdm")
cdm

cdm |>
   cdmSelect("person")
```

cdmSource

Get the cdmSource of an object.

Description

Get the cdmSource of an object.

Usage

```
cdmSource(x, cdm = lifecycle::deprecated())
```

Arguments

x Object to obtain the cdmSource.

cdm Deprecated, use x please.

Value

A cdm_source object.

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
cdm <- cdmFromTables(</pre>
  tables = list(
    "person" = tibble(
      person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
      race_concept_id = 0, ethnicity_concept_id = 0
   ),
    "observation_period" = tibble(
      observation_period_id = 1:3, person_id = 1:3,
      observation_period_start_date = as.Date("2000-01-01"),
      observation_period_end_date = as.Date("2023-12-31"),
      period_type_concept_id = 0
   )
  ),
  cdmName = "mock"
)
```

24 cdmSourceType

```
cdmSource(cdm)
cdmSource(cdm$person)
```

cdmSourceType

Get the source type of a cdm_reference object.

Description

[Deprecated]

Usage

```
cdmSourceType(cdm)
```

Arguments

cdm

A cdm_reference object.

Value

A character vector with the type of source of the cdm_reference object.

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
cdm <- cdmFromTables(</pre>
  tables = list(
    "person" = tibble(
      person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
      race_concept_id = 0, ethnicity_concept_id = 0
   ),
    "observation_period" = tibble(
      observation_period_id = 1:3, person_id = 1:3,
      observation_period_start_date = as.Date("2000-01-01"),
      observation_period_end_date = as.Date("2023-12-31"),
      period_type_concept_id = 0
   )
  ),
  cdmName = "mock"
)
cdmSourceType(cdm)
```

cdmTableFromSource 25

cdmTableFromSource	This is an internal developer focused function that creates a cdm table
Culli labitel i ollisodi ce	1 3 3 =
	from a table that shares the source but it is not a cdm_table. Please
	use insertTable if you want to insert a table to a cdm reference object.

Description

This is an internal developer focused function that creates a cdm_table from a table that shares the source but it is not a cdm_table. Please use insertTable if you want to insert a table to a cdm_reference object.

Usage

```
cdmTableFromSource(src, value)
```

Arguments

src A cdm_source object.

value A table that shares source with the cdm_reference object.

Value

A cdm_table.

cdmVersion

Get the version of an object.

Description

Get the version of an object.

Usage

cdmVersion(x)

Arguments

X

Object to know the cdm version of an object.

Value

A character vector indicating the cdm version.

Examples

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
cdm <- cdmFromTables(</pre>
  tables = list(
    "person" = tibble(
      person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
      race_concept_id = 0, ethnicity_concept_id = 0
    "observation_period" = tibble(
      observation_period_id = 1:3, person_id = 1:3,
      observation_period_start_date = as.Date("2000-01-01"),
      observation_period_end_date = as.Date("2023-12-31"),
      period_type_concept_id = 0
   )
  ),
  cdmName = "mock"
)
cdmVersion(cdm)
cdmVersion(cdm$person)
```

checkCohortRequirements

Check whether a cohort table satisfies requirements

Description

[Deprecated]

Usage

```
checkCohortRequirements(
  cohort,
  checkEndAfterStart = TRUE,
  checkOverlappingEntries = TRUE,
  checkMissingValues = TRUE,
  checkInObservation = TRUE,
  type = "error",
  call = parent.frame()
)
```

Arguments

cohort cohort_table object.

cohortCodelist 27

checkEndAfterStart

If TRUE a check that all cohort end dates come on or after cohort start date will be performed.

checkOverlappingEntries

If TRUE a check that no individuals have overlapping cohort entries will be performed.

checkMissingValues

If TRUE a check that there are no missing values in required fields will be performed

checkInObservation

If TRUE a check that cohort entries are within the individuals observation periods will be performed.

ods will be performed

Can be either "error" or "warning". If "error" any check failure will result in an

error, whereas if "warning" any check failure will result in a warning.

call The call for which to return the error message.

Value

type

An error will be returned if any of the selected checks fail.

cohortCodelist

Get codelist from a cohort_table object.

Description

Get codelist from a cohort_table object.

Usage

```
cohortCodelist(
  cohortTable,
  cohortId,
  type = c("index event", "inclusion criteria", "exclusion criteria", "exit criteria")
)
```

Arguments

cohortTable A cohort_table object.

cohortId A particular cohort definition id that is present in the cohort table.

type The reason for the codelist. Can be "index event", "inclusion criteria", or "exit

criteria".

Value

A table with the codelists used.

28 cohortColumns

Examples

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
person <- tibble(</pre>
 person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
 race_concept_id = 0, ethnicity_concept_id = 0
observation_period <- tibble(</pre>
 observation_period_id = 1, person_id = 1,
 observation_period_start_date = as.Date("2000-01-01"),
 observation_period_end_date = as.Date("2023-12-31"),
 period_type_concept_id = 0
)
cohort <- tibble(</pre>
 cohort_definition_id = c(1, 1, 1, 2),
 subject_id = 1,
 cohort_start_date = as.Date(c(
    "2020-01-01", "2021-01-01", "2022-01-01", "2022-01-01"
 )),
 cohort_end_date = as.Date(c(
    "2020-01-01", "2021-01-01", "2022-01-01", "2022-01-01"
 ))
)
cdm <- cdmFromTables(</pre>
 tables = list("person" = person, "observation_period" = observation_period),
 cdmName = "my_example_cdm",
 cohortTables = list("cohort1" = cohort)
)
cdm$cohort1 <- newCohortTable(table = cdm$cohort1,</pre>
                                 cohortCodelistRef = dplyr::tibble(
                                 cohort_definition_id = c(1,1,1,2,2),
                                 codelist_name =c("disease X", "disease X", "disease X",
                                                   "disease Y", "disease Y"),
                                 concept_id = c(1,2,3,4,5),
                                 type = "index event"
cohortCodelist(cdm$cohort1, cohortId = 1, type = "index event")
```

cohortColumns

Required columns for a generated cohort set.

Description

Required columns for a generated cohort set.

Usage

```
cohortColumns(table, version = "5.3")
```

cohortCount 29

Arguments

table Either cohort, cohort_set or cohort_attrition version Version of the OMOP Common Data Model.

Value

Character vector with the column names Required columns

Examples

```
library(omopgenerics)
cohortColumns("cohort")
```

cohortCount

Get cohort counts from a cohort_table object.

Description

Get cohort counts from a cohort_table object.

Usage

```
cohortCount(cohort)
```

Arguments

cohort

A cohort_table object.

Value

A table with the counts.

30 cohortTables

```
cohort <- tibble(
  cohort_definition_id = c(1, 1, 1, 2),
  subject_id = 1,
  cohort_start_date = as.Date(c(
      "2020-01-01", "2021-01-01", "2022-01-01", "2022-01-01"
)),
  cohort_end_date = as.Date(c(
      "2020-01-01", "2021-01-01", "2022-01-01", "2022-01-01"
)),
))
cdm <- cdmFromTables(
  tables = list("person" = person, "observation_period" = observation_period),
  cdmName = "my_example_cdm",
  cohortTables = list("cohort1" = cohort)
)
cohortCount(cdm$cohort1)</pre>
```

cohortTables

Cohort tables that a cdm reference can contain in the OMOP Common Data Model.

Description

Cohort tables that a cdm reference can contain in the OMOP Common Data Model.

Usage

```
cohortTables(version = "5.3")
```

Arguments

version

Version of the OMOP Common Data Model.

Value

cohort tables

```
library(omopgenerics)
cohortTables()
```

collect.cdm_reference 31

collect.cdm_reference Retrieves the cdm reference into a local cdm.

Description

Retrieves the cdm reference into a local cdm.

Usage

```
## S3 method for class 'cdm_reference'
collect(x, ...)
```

Arguments

x A cdm_reference object.

... For compatibility only, not used.

Value

A local cdm_reference.

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
cdm <- cdmFromTables(</pre>
  tables = list(
    "person" = dplyr::tibble(
      person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
      race_concept_id = 0, ethnicity_concept_id = 0
    "observation_period" = dplyr::tibble(
      observation_period_id = 1:3, person_id = 1:3,
      observation_period_start_date = as.Date("2000-01-01"),
      observation_period_end_date = as.Date("2023-12-31"),
      period_type_concept_id = 0
   )
  ),
  cdmName = "mock"
collect(cdm)
```

32 combineStrata

```
collect.cohort_table To collect a cohort_table object.
```

Description

To collect a cohort_table object.

Usage

```
## S3 method for class 'cohort_table'
collect(x, ...)
```

Arguments

x cohort_table object.

... Not used (for compatibility).

Value

A data frame with the cohort_table

combineStrata

Provide all combinations of strata levels.

Description

Provide all combinations of strata levels.

Usage

```
combineStrata(levels)
```

Arguments

levels

Vector of all strata levels to combine.

Value

A vector of all combinations of strata.

compute.cdm_table 33

compute.cdm_table

Store results in a table.

Description

Store results in a table.

Usage

```
## $3 method for class 'cdm_table'
compute(
    x,
    name = NULL,
    temporary = NULL,
    overwrite = TRUE,
    logPrefix = NULL,
    ...
)
```

Arguments

x Table in the cdm.

name Name to store the table with.

temporary Whether to store table temporarily (TRUE) or permanently (FALSE).

overwrite Whether to overwrite previously existing table with name same.

logPrefix Prefix to use when saving a log file.

... For compatibility (not used).

Value

Reference to a table in the cdm

dropSourceTable

Drop a table from a cdm object.

Description

Drop a table from a cdm object.

Usage

```
dropSourceTable(cdm, name)
```

34 dropTable

Arguments

cdm A cdm reference.

name Name(s) of the table(s) to insert. Tidyselect statements are supported.

Value

The table in the cdm reference.

dropTable

Drop a table from a cdm object.

Description

Drop a table from a cdm object.

Usage

```
dropTable(cdm, name)
```

Arguments

cdm A cdm reference.

name Name(s) of the table(s) to drop Tidyselect statements are supported.

Value

The cdm reference.

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
person <- tibble(</pre>
  person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
  race_concept_id = 0, ethnicity_concept_id = 0
)
observation_period <- tibble(</pre>
  observation_period_id = 1, person_id = 1,
  observation_period_start_date = as.Date("2000-01-01"),
  observation_period_end_date = as.Date("2023-12-31"),
  period_type_concept_id = 0
cohort <- tibble(</pre>
  cohort_definition_id = c(1, 1, 1, 2),
  subject_id = 1,
  cohort_start_date = as.Date(c(
    "2020-01-01", "2021-01-01", "2022-01-01", "2022-01-01"
```

emptyAchillesTable 35

```
)),
  cohort_end_date = as.Date(c(
    "2020-01-01", "2021-01-01", "2022-01-01", "2022-01-01"
)),
)

cdm <- cdmFromTables(
  tables = list("person" = person, "observation_period" = observation_period),
  cdmName = "my_example_cdm",
  cohortTables = list("cohort1" = cohort)
)

cdm

cdm <- dropTable(cdm = cdm, name = "cohort1")

cdm</pre>
```

 ${\tt emptyAchillesTable}$

Create an empty achilles table

Description

Create an empty achilles table

Usage

```
emptyAchillesTable(cdm, name)
```

Arguments

cdm A cdm_reference to create the table.

name Name of the table to create.

Value

The cdm_reference with an achilles empty table

```
library(omopgenerics)
cdm <- emptyCdmReference("my_example_cdm")
emptyAchillesTable(cdm = cdm, name = "achilles_results")</pre>
```

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 ${\tt emptyCdmReference}$

Create an empty cdm_reference

Description

Create an empty cdm_reference

Usage

```
emptyCdmReference(cdmName, cdmVersion = NULL)
```

Arguments

cdmName Name of the cdm_reference cdmVersion Version of the cdm_reference

Value

An empty cdm_reference

Examples

```
library(omopgenerics)
emptyCdmReference(cdmName = "my_example_cdm")
```

emptyCodelist

Empty codelist object.

Description

Empty codelist object.

Usage

```
emptyCodelist()
```

Value

An empty codelist object.

```
emptyCodelist()
```

emptyCodelistWithDetails

Empty codelist object.

Description

Empty codelist object.

Usage

```
emptyCodelistWithDetails()
```

Value

An empty codelist object.

Examples

```
emptyCodelistWithDetails()
```

 ${\tt emptyCohortTable}$

Create an empty cohort_table object

Description

Create an empty cohort_table object

Usage

```
emptyCohortTable(cdm, name, overwrite = TRUE)
```

Arguments

cdm A cdm_reference to create the table.

name Name of the table to create.

overwrite Whether to overwrite an existent table.

Value

The cdm_reference with an empty cohort table

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Examples

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
person <- tibble(</pre>
  person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
  race_concept_id = 0, ethnicity_concept_id = 0
observation_period <- tibble(</pre>
  observation_period_id = 1, person_id = 1,
  observation_period_start_date = as.Date("2000-01-01"),
  observation_period_end_date = as.Date("2023-12-31"),
  period_type_concept_id = 0
)
cdm <- cdmFromTables(</pre>
  tables = list("person" = person, "observation_period" = observation_period),
  cdmName = "test"
)
cdm <- emptyCohortTable(cdm, "my_empty_cohort")</pre>
cdm
cdm$my_empty_cohort
settings(cdm$my_empty_cohort)
attrition(cdm$my_empty_cohort)
cohortCount(cdm$my_empty_cohort)
```

emptyOmopTable

Create an empty omop table

Description

Create an empty omop table

Usage

```
emptyOmopTable(cdm, name)
```

Arguments

cdm A cdm_reference to create the table.

name Name of the table to create.

Value

The cdm_reference with an empty cohort table

Examples

```
library(omopgenerics)
person <- dplyr::tibble(</pre>
  person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
  race_concept_id = 0, ethnicity_concept_id = 0
)
observation_period <- dplyr::tibble(</pre>
  observation_period_id = 1, person_id = 1,
  observation_period_start_date = as.Date("2000-01-01"),
  observation_period_end_date = as.Date("2023-12-31"),
  period_type_concept_id = 0
)
cdm <- cdmFromTables(</pre>
  tables = list("person" = person, "observation_period" = observation_period),
  cdmName = "test"
cdm <- emptyOmopTable(cdm, "drug_exposure")</pre>
cdm$drug_exposure
```

emptySummarisedResult Empty summarised_result object.

Description

Empty summarised_result object.

Usage

```
emptySummarisedResult(settings = NULL)
```

Arguments

settings

Tibble/data.frame with the settings of the empty summarised_result. It has to contain at least result_id column.

Value

An empty summarised_result object.

```
library(omopgenerics)
emptySummarisedResult()
```

40 exportCodelist

estimate Type Choices

Choices that can be present in estimate_type column.

Description

Choices that can be present in estimate_type column.

Usage

```
estimateTypeChoices()
```

Value

A character vector with the options that can be present in estimate_type column in the summarised_result objects.

Examples

```
library(omopgenerics)
estimateTypeChoices()
```

exportCodelist

Export a codelist object.

Description

Export a codelist object.

Usage

```
exportCodelist(x, path, type = "json")
```

Arguments

x A codelist

path Path to where files will be created.

type Type of files to export. Currently 'json' and 'csv' are supported.

Value

Files with codelists

 ${\tt exportConceptSetExpression}$

Export a concept set expression.

Description

Export a concept set expression.

Usage

```
exportConceptSetExpression(x, path, type = "json")
```

Arguments

x A concept set expression

path Path to where files will be created.

type Type of files to export. Currently 'json' and 'csv' are supported.

Value

Files with concept set expressions

```
exportSummarisedResult
```

Export a summarised_result object to a csv file.

Description

Export a summarised_result object to a csv file.

Usage

```
exportSummarisedResult(
    ...,
    minCellCount = 5,
    fileName = "results_{cdm_name}_{date}.csv",
    path = getwd()
)
```

42 filterAdditional

Arguments

... A set of summarised_result objects.

minCellCount Minimum count for suppression purposes.

fileName Name of the file that will be created. Use {cdm_name} to refer to the cdmName

of the objects and {date} to add the export date.

path Path where to create the csv file. It is ignored if fileName it is a full name with

path included.

 $filter \verb|Additional| filter the additional_name-additional_level pair in a sum-$

marised_result

Description

Filter the additional_name-additional_level pair in a summarised_result

Usage

```
filterAdditional(result, ...)
```

Arguments

result A <summarised_result> object.

Expressions that return a logical value (additionalColumns() are used to evaluate the expression and are defined in terms of the exprisible in data. If multiple

uate the expression), and are defined in terms of the variables in .data. If multiple expressions are included, they are combined with the & operator. Only rows for

which all conditions evaluate to TRUE are kept.

Value

A <summarised_result> object with only the rows that fulfill the required specified additional.

```
library(dplyr)
library(omopgenerics)

x <- tibble(
   "result_id" = 1L,
   "cdm_name" = "eunomia",
   "group_name" = "cohort_name",
   "group_level" = c("cohort1", "cohort2", "cohort3"),
   "strata_name" = "sex",
   "strata_level" = "Female",
   "variable_name" = "number subjects",
   "variable_level" = NA_character_,
   "estimate_name" = "count",</pre>
```

filterGroup 43

```
"estimate_type" = "integer",
"estimate_value" = c("100", "44", "14"),
"additional_name" = c("year", "time_step", "year &&& time_step"),
"additional_level" = c("2010", "4", "2015 &&& 5")
) |>
newSummarisedResult()

x |>
filterAdditional(year == "2010")
```

filterGroup

Filter the group_name-group_level pair in a summarised_result

Description

Filter the group_name-group_level pair in a summarised_result

Usage

```
filterGroup(result, ...)
```

Arguments

result

A <summarised_result> object.

. . .

Expressions that return a logical value (groupColumns() are used to evaluate the expression), and are defined in terms of the variables in .data. If multiple expressions are included, they are combined with the & operator. Only rows for which all conditions evaluate to TRUE are kept.

Value

A <summarised_result> object with only the rows that fulfill the required specified group.

```
library(dplyr)
library(omopgenerics)

x <- tibble(
    "result_id" = 1L,
    "cdm_name" = "eunomia",
    "group_name" = c("cohort_name", "age_group &&& cohort_name", "age_group"),
    "group_level" = c("my_cohort", ">40 &&& second_cohort", "<40"),
    "strata_name" = "sex",
    "strata_level" = "Female",
    "variable_name" = "number subjects",
    "variable_level" = NA_character_,
    "estimate_name" = "count",</pre>
```

44 filterSettings

```
"estimate_type" = "integer",
  "estimate_value" = c("100", "44", "14"),
  "additional_name" = "overall",
  "additional_level" = "overall"
) |>
  newSummarisedResult()

x |>
  filterGroup(cohort_name == "second_cohort")
```

filterSettings

Filter a <summarised_result> using the settings

Description

Filter a <summarised_result> using the settings

Usage

```
filterSettings(result, ...)
```

Arguments

result

A <summarised_result> object.

. . .

Expressions that return a logical value (columns in settings are used to evaluate the expression), and are defined in terms of the variables in .data. If multiple expressions are included, they are combined with the & operator. Only rows for which all conditions evaluate to TRUE are kept.

Value

A <summarised_result> object with only the result_id rows that fulfill the required specified settings.

```
library(dplyr)
library(omopgenerics)

x <- tibble(
   "result_id" = as.integer(c(1, 2)),
   "cdm_name" = c("cprd", "eunomia"),
   "group_name" = "cohort_name",
   "group_level" = "my_cohort",
   "strata_name" = "sex",
   "strata_level" = "male",
   "variable_name" = "Age group",
   "variable_level" = "10 to 50",</pre>
```

filterStrata 45

```
"estimate_name" = "count",
"estimate_type" = "numeric",
"estimate_value" = "5",
"additional_name" = "overall",
"additional_level" = "overall"
) |>
  newSummarisedResult(settings = tibble(
    "result_id" = c(1, 2), "custom" = c("A", "B")
))

x

x |> filterSettings(custom == "A")
```

filterStrata

Filter the strata_name-strata_level pair in a summarised_result

Description

Filter the strata_name-strata_level pair in a summarised_result

Usage

```
filterStrata(result, ...)
```

Arguments

result

A <summarised_result> object.

Expressions that return a logical value (strataColumns() are used to evaluate the expression), and are defined in terms of the variables in .data. If multiple expressions are included, they are combined with the & operator. Only rows for which all conditions evaluate to TRUE are kept.

Value

A <summarised_result> object with only the rows that fulfill the required specified strata.

```
library(dplyr)
library(omopgenerics)

x <- tibble(
   "result_id" = 1L,
   "cdm_name" = "eunomia",
   "group_name" = "cohort_name",
   "group_level" = "my_cohort",
   "strata_name" = c("sex", "sex &&& age_group", "sex &&& year"),</pre>
```

46 getCohortName

```
"strata_level" = c("Female", "Male &&& <40", "Female &&& 2010"),
"variable_name" = "number subjects",
"variable_level" = NA_character_,
"estimate_name" = "count",
"estimate_type" = "integer",
"estimate_value" = c("100", "44", "14"),
"additional_name" = "overall",
"additional_level" = "overall"
) |>
newSummarisedResult()

x |>
filterStrata(sex == "Female")
```

getCohortId

Get the cohort definition id of a certain name

Description

Get the cohort definition id of a certain name

Usage

```
getCohortId(cohort, cohortName = NULL)
```

Arguments

cohort A cohort_table object.

cohortName Names of the cohort of interest. If NULL all cohort names are shown.

Value

Cohort definition ids

getCohortName

Get the cohort name of a certain cohort definition id

Description

Get the cohort name of a certain cohort definition id

Usage

```
getCohortName(cohort, cohortId = NULL)
```

getPersonIdentifier 47

Arguments

cohort A cohort_table object.

cohortId Cohort definition id of interest. If NULL all cohort ids are shown.

Value

Cohort names

 ${\tt getPersonIdentifier} \qquad \textit{Get the column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a table (either a column name with the person identifier from a column name with the person identifier from a table (either a column name with the person identifier from a column name with the$

subject_id or person_id), it will throw an error if it contains both or

neither.

Description

Get the column name with the person identifier from a table (either subject_id or person_id), it will throw an error if it contains both or neither.

Usage

```
getPersonIdentifier(x, call = parent.frame())
```

Arguments

x A table.

call A call argument passed to cli functions.

Value

Person identifier column.

Description

Identifies and returns the unique values in group_name column.

Usage

```
groupColumns(result)
```

Arguments

result A tibble.

48 importCodelist

Value

Unique values of the group name column.

Examples

```
{
 library(dplyr)
 library(omopgenerics)
 x <- tibble(
   "result_id" = as.integer(c(1, 2)),
    "cdm_name" = c("cprd", "eunomia"),
    "group_name" = "cohort",
    "group_level" = "my_cohort",
    "strata_name" = "sex",
    "strata_level" = "male",
    "variable_name" = "Age group",
    "variable_level" = "10 to 50",
    "estimate_name" = "count",
    "estimate_type" = "numeric",
    "estimate_value" = "5",
    "additional_name" = "overall",
    "additional_level" = "overall"
 ) |>
    newSummarisedResult(settings = tibble(
      "result_id" = c(1, 2), "custom" = c("A", "B")
   ))
 x |> groupColumns()
```

importCodelist

Import a codelist.

Description

Import a codelist.

Usage

```
importCodelist(path, type = "json")
```

Arguments

path Path to where files will be created.

type Type of files to export. Currently 'json' and 'csv' are supported.

Value

A codelist

import Concept Set Expression

Import a concept set expression.

Description

Import a concept set expression.

Usage

```
importConceptSetExpression(path, type = "json")
```

Arguments

path Path to where files will be created.

type Type of files to export. Currently 'json' and 'csv' are supported.

Value

A concept set expression

importSummarisedResult

Import a set of summarised results.

Description

Import a set of summarised results.

Usage

```
importSummarisedResult(path, recursive = FALSE)
```

Arguments

path Path to directory with CSV files containing summarised results or to a specific

CSV file with a summarised result.

recursive If TRUE and path is a directory, search for files will recurse into directories

Value

A summarised result

50 insertTable

insertFromSource

Convert a table that is not a cdm_table but have the same original source to a cdm_table. This Table is not meant to be used to insert tables in the cdm, please use insertTable instead.

Description

[Deprecated]

Usage

```
insertFromSource(cdm, value)
```

Arguments

cdm A cdm_reference object.

value A table that shares source with the cdm_reference object.

Value

A table in the cdm_reference environment

insertTable

Insert a table to a cdm object.

Description

Insert a table to a cdm object.

Usage

```
insertTable(cdm, name, table, overwrite = TRUE, temporary = FALSE)
```

Arguments

cdm A cdm reference or the source of a cdm reference.

name Name of the table to insert.
table Table to insert to the cdm.

overwrite Whether to overwrite an existent table. temporary Whether to create a temporary table.

isResultSuppressed 51

Value

```
The cdm reference. 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("2023-12-31"), period_type_concept_id = 0 ) cdm <- cdmFromTables( tables = list("person" = person, "observation_period" = observation_period), cdmName = "my_example_cdm" ) x <- tibble(a = 1)

cdm <- insertTable(cdm = cdm, name = "new_table", table = x)

cdm$new_table
```

isResultSuppressed

isResultSuppressed

Description

isResultSuppressed

Usage

```
isResultSuppressed(result, minCellCount = 5)
```

Arguments

result The suppressed result to check

minCellCount Minimum count of records used when suppressing

Value

Warning or message with check result

isTableEmpty

Check if a table is empty or not

Description

Check if a table is empty or not

Usage

```
isTableEmpty(table)
```

Arguments

table

a table

52 newAchillesTable

Value

Boolean to indicate if a cdm_table is empty (TRUE or FALSE).

listSourceTables

List tables that can be accessed though a cdm object.

Description

List tables that can be accessed though a cdm object.

Usage

```
listSourceTables(cdm)
```

Arguments

cdm

A cdm reference or the source of a cdm reference.

Value

A character vector with the names of tables.

newAchillesTable

Create an achilles table from a cdm_table.

Description

Create an achilles table from a cdm_table.

Usage

```
newAchillesTable(table, version = "5.3", cast = FALSE)
```

Arguments

table A cdm_table.

version version of the cdm.

cast Whether to cast columns to the correct type.

Value

An achilles_table object

newCdmReference 53

newCdmReference cdm_reference objects constructor

Description

cdm_reference objects constructor

Usage

```
newCdmReference(tables, cdmName, cdmVersion = NULL, .softValidation = FALSE)
```

Arguments

tables List of tables that are part of the OMOP Common Data Model reference.

cdmName Name of the cdm object.

cdmVersion Version of the cdm. Supported versions 5.3 and 5.4.

.softValidation

Whether to perform a soft validation of consistency. If set to FALSE, non overlapping observation periods are ensured.

Value

A cdm_reference object.

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
cdmTables <- list(</pre>
  "person" = tibble(
   person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
   race_concept_id = 0, ethnicity_concept_id = 0
   newCdmTable(newLocalSource(), "person"),
  "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("2023-12-31"),
    period_type_concept_id = 0
 ) |>
    newCdmTable(newLocalSource(), "observation_period")
)
cdm <- newCdmReference(tables = cdmTables, cdmName = "mock")</pre>
cdm
```

54 newCdmTable

newCdmSource

Create a cdm source object.

Description

Create a cdm source object.

Usage

```
newCdmSource(src, sourceType)
```

Arguments

src Source to a cdm object. sourceType Type of the source object.

Value

A validated cdm source object.

newCdmTable

Create an cdm table.

Description

Create an cdm table.

Usage

```
newCdmTable(table, src, name)
```

Arguments

table A table that is part of a cdm.
src The source of the table.
name The name of the table.

Value

A cdm_table object

newCodelist 55

newCodelist

'codelist' object constructor

Description

'codelist' object constructor

Usage

```
newCodelist(x)
```

Arguments

Х

A named list where each element contains a vector of concept IDs.

Value

A codelist object.

newCodelistWithDetails

'codelist' object constructor

Description

'codelist' object constructor

Usage

```
newCodelistWithDetails(x)
```

Arguments

Х

A named list where each element contains a tibble with the column concept_id

Value

A codelist object.

56 newCohortTable

newCohortTable

cohort_table objects constructor.

Description

cohort_table objects constructor.

Usage

```
newCohortTable(
  table,
  cohortSetRef = attr(table, "cohort_set"),
  cohortAttritionRef = attr(table, "cohort_attrition"),
  cohortCodelistRef = attr(table, "cohort_codelist"),
    .softValidation = FALSE
)
```

Arguments

table

cdm_table object with at least: cohort_definition_id, subject_id, cohort_start_date, cohort_end_date.

cohortSetRef

Table with at least: cohort_definition_id, cohort_name

cohortAttritionRef

Table with at least: cohort_definition_id, number_subjects, number_records, reason_id, reason, excluded_subjects, excluded_records.

cohortCodelistRef

Table with at least: cohort_definition_id, codelist_name, and concept_id.

.softValidation

Whether to perform a soft validation of consistency. If set to FALSE four additional checks will be performed: 1) a check that cohort end date is not before cohort start date, 2) a check that there are no missing values in required columns, 3) a check that cohort duration is all within observation period, and 4) that there are no overlapping cohort entries

Value

A cohort_table object

```
person <- dplyr::tibble(
   person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
   race_concept_id = 0, ethnicity_concept_id = 0
)
observation_period <- dplyr::tibble(
   observation_period_id = 1, person_id = 1,
   observation_period_start_date = as.Date("2000-01-01"),</pre>
```

```
observation_period_end_date = as.Date("2023-12-31"),
  period_type_concept_id = 0
)
cohort1 <- dplyr::tibble(</pre>
  cohort_definition_id = 1, subject_id = 1,
  cohort_start_date = as.Date("2020-01-01"),
  cohort_end_date = as.Date("2020-01-10")
cdm <- cdmFromTables(</pre>
  tables = list(
    "person" = person,
    "observation_period" = observation_period,
    "cohort1" = cohort1
  ),
  cdmName = "test"
)
\mathsf{cdm}
cdm$cohort1 <- newCohortTable(table = cdm$cohort1)</pre>
settings(cdm$cohort1)
attrition(cdm$cohort1)
cohortCount(cdm$cohort1)
```

 ${\tt newConceptSetExpression}$

'conceptSetExpression' object constructor

Description

'conceptSetExpression' object constructor

Usage

```
newConceptSetExpression(x)
```

Arguments

Х

a named list of tibbles, each of which containing concept set definitions

Value

A conceptSetExpression

58 newOmopTable

 ${\tt newLocalSource}$

A new local source for the cdm

Description

A new local source for the cdm

Usage

```
newLocalSource()
```

Value

A list in the format of a cdm source

Examples

```
library(omopgenerics)
newLocalSource()
```

newOmopTable

Create an omop table from a cdm table.

Description

Create an omop table from a cdm table.

Usage

```
newOmopTable(table, version = "5.3", cast = FALSE)
```

Arguments

table A cdm_table.

version version of the cdm.

cast Whether to cast columns to the correct type.

Value

An omop_table object

newSummarisedResult 59

newSummarisedResult 'summarised_results' object constructor

Description

'summarised_results' object constructor

Usage

```
newSummarisedResult(x, settings = attr(x, "settings"))
```

Arguments

x Table.

settings Settings for the summarised_result object.

Value

A summarised_result object

```
library(dplyr)
library(omopgenerics)
x <- tibble(</pre>
  "result_id" = 1L,
  "cdm_name" = "cprd",
  "group_name" = "cohort_name",
  "group_level" = "acetaminophen",
  "strata_name" = "sex &&& age_group",
"strata_level" = c("male &&& <40", "male &&& >=40"),
  "variable_name" = "number_subjects",
  "variable_level" = NA_character_,
  "estimate_name" = "count",
  "estimate_type" = "integer",
  "estimate_value" = c("5", "15"),
  "additional_name" = "overall",
  "additional_level" = "overall"
) |>
  newSummarisedResult()
settings(x)
summary(x)
x <- tibble(</pre>
  "result_id" = 1L,
  "cdm_name" = "cprd",
```

60 omopColumns

```
"group_name" = "cohort_name",
  "group_level" = "acetaminophen",
  "strata_name" = "sex &&& age_group",
  "strata_level" = c("male &&& <40", "male &&& >=40"),
  "variable_name" = "number_subjects",
  "variable_level" = NA_character_,
  "estimate_name" = "count",
  "estimate_type" = "integer",
  "estimate_value" = c("5", "15"),
  "additional_name" = "overall",
  "additional_level" = "overall"
) |>
 newSummarisedResult(settings = tibble(
    result_id = 1L, result_type = "custom_summary", mock = TRUE, value = 5
 ))
Х
settings(x)
summary(x)
```

omopColumns

Required columns that the standard tables in the OMOP Common Data Model must have.

Description

Required columns that the standard tables in the OMOP Common Data Model must have.

Usage

```
omopColumns(
  table,
  field = NULL,
  version = "5.3",
  onlyRequired = lifecycle::deprecated()
)
```

Arguments

table Table to see required columns.

field Name of the specific field.

version Version of the OMOP Common Data Model.

onlyRequired deprecated

Value

Character vector with the column names

omopTableFields 61

Examples

```
library(omopgenerics)
omopColumns("person")
```

omopTableFields

Return a table of omop cdm fields informations

Description

Return a table of omop cdm fields informations

Usage

```
omopTableFields(cdmVersion = "5.3")
```

Arguments

cdmVersion

cdm version of the omop cdm.

Value

a tibble contain informations on all the different fields in omop cdm.

omopTables

Standard tables that a cdm reference can contain in the OMOP Common Data Model.

Description

Standard tables that a cdm reference can contain in the OMOP Common Data Model.

Usage

```
omopTables(version = "5.3")
```

Arguments

version

Version of the OMOP Common Data Model.

Value

Standard tables

62 pivotEstimates

Examples

```
library(omopgenerics)
omopTables()
```

pivotEstimates

Set estimates as columns

Description

Pivot the estimates as new columns in result table.

Usage

```
pivotEstimates(result, pivotEstimatesBy = "estimate_name", nameStyle = NULL)
```

Arguments

Names from which pivot wider the estimate values. If NULL the table will not be pivotted.

nameStyle

Name style (glue package specifications) to customise names when pivotting estimates. If NULL standard tidyr::pivot_wider formatting will be used.

Value

A tibble.

```
library(dplyr)
library(omopgenerics)

x <- tibble(
   "result_id" = 1L,
   "cdm_name" = c("cprd", "eunomia"),
   "group_name" = "cohort_name",
   "group_level" = "my_cohort",
   "strata_name" = "sex",
   "strata_level" = "male",
   "variable_name" = "Age group",
   "variable_level" = "10 to 50",
   "estimate_name" = "count",
   "estimate_type" = "numeric",
   "estimate_value" = "5",</pre>
```

print.cdm_reference 63

```
"additional_name" = "overall",
    "additional_level" = "overall"
) |>
    newSummarisedResult()

x |>
    pivotEstimates()
}
```

print.cdm_reference

Print a CDM reference object

Description

Print a CDM reference object

Usage

```
## S3 method for class 'cdm_reference'
print(x, ...)
```

Arguments

x A cdm_reference object

... Included for compatibility with generic. Not used.

Value

Invisibly returns the input

```
library(omopgenerics)

cdm <- cdmFromTables(
   tables = list(
    "person" = dplyr::tibble(
        person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
        race_concept_id = 0, ethnicity_concept_id = 0
   ),
    "observation_period" = dplyr::tibble(
        observation_period_id = 1:3, person_id = 1:3,
        observation_period_start_date = as.Date("2000-01-01"),
        observation_period_end_date = as.Date("2023-12-31"),
        period_type_concept_id = 0
   )
   ),
   cdmName = "mock"</pre>
```

```
)
print(cdm)
```

print.codelist

Print a codelist

Description

Print a codelist

Usage

```
## S3 method for class 'codelist'
print(x, ...)
```

Arguments

x A codelist

... Included for compatibility with generic. Not used.

Value

Invisibly returns the input

Examples

```
codes <- list("disease X" = c(1, 2, 3), "disease Y" = c(4, 5)) codes <- newCodelist(codes) print(codes)
```

```
print.codelist_with_details
```

Print a codelist with details

Description

Print a codelist with details

Usage

```
## S3 method for class 'codelist_with_details'
print(x, ...)
```

Arguments

- x A codelist with details
- ... Included for compatibility with generic. Not used.

Value

Invisibly returns the input

Examples

```
codes <- list("disease X" = dplyr::tibble(
  concept_id = c(1, 2, 3),
  other = c("a", "b", "c")
))
codes <- newCodelistWithDetails(codes)
print(codes)</pre>
```

print.conceptSetExpression

Print a concept set expression

Description

Print a concept set expression

Usage

```
## S3 method for class 'conceptSetExpression' print(x, ...)
```

Arguments

- x A concept set expression
- ... Included for compatibility with generic. Not used.

Value

Invisibly returns the input

66 readSourceTable

Examples

```
asthma_cs <- list(
  "asthma_narrow" = dplyr::tibble(
  "concept_id" = 1,
  "excluded" = FALSE,
  "descendants" = TRUE,
  "mapped" = FALSE
),
  "asthma_broad" = dplyr::tibble(
  "concept_id" = c(1, 2),
  "excluded" = FALSE,
  "descendants" = TRUE,
  "mapped" = FALSE
)
)
asthma_cs <- newConceptSetExpression(asthma_cs)
print(asthma_cs)</pre>
```

readSourceTable

Read a table from the cdm_source and add it to to the cdm.

Description

Read a table from the cdm_source and add it to to the cdm.

Usage

```
readSourceTable(cdm, name)
```

Arguments

cdm A cdm reference.

name Name of a table to read in the cdm_source space.

Value

A cdm_reference with new table.

recordCohortAttrition 67

recordCohortAttrition Update cohort attrition.

Description

Update cohort attrition.

Usage

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

Arguments

cohort A cohort_table object.
reason A character string.

cohortId Cohort definition id of the cohort to update attrition. If NULL all cohort_definition_id

are updated.

Value

cohort_table with updated attrition.

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
person <- tibble(</pre>
  person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
  race_concept_id = 0, ethnicity_concept_id = 0
observation_period <- tibble(</pre>
  observation_period_id = 1, person_id = 1,
  observation_period_start_date = as.Date("2000-01-01"),
  observation_period_end_date = as.Date("2023-12-31"),
  period_type_concept_id = 0
)
cohort <- tibble(</pre>
  cohort_definition_id = c(1, 1, 1, 2),
  subject_id = 1,
 cohort_start_date = as.Date(c("2020-01-01", "2021-01-01", "2022-01-01", "2022-01-01")),
  cohort_end_date = as.Date(c("2020-01-01", "2021-01-01", "2022-01-01", "2022-01-01")),
cdm <- cdmFromTables(</pre>
  tables = list("person" = person, "observation_period" = observation_period),
  cdmName = "my_example_cdm",
  cohortTables = list("cohort1" = cohort)
)
```

68 resultColumns

```
cdm$cohort1
attrition(cdm$cohort1)

cdm$cohort1 <- cdm$cohort1 |>
    group_by(cohort_definition_id, subject_id) |>
    filter(cohort_start_date == min(cohort_start_date)) |>
    ungroup() |>
    compute(name = "cohort1", temporary = FALSE) |>
    recordCohortAttrition("Restrict to first observation")

cdm$cohort1
attrition(cdm$cohort1)
```

resultColumns

Required columns that the result tables must have.

Description

Required columns that the result tables must have.

Usage

```
resultColumns(table = "summarised_result")
```

Arguments

table

Table to see required columns.

Value

Required columns

```
library(omopgenerics)
resultColumns()
```

resultPackageVersion 69

resultPackageVersion

Check if different packages version are used for summarise_results object

Description

Check if different packages version are used for summarise_results object

Usage

```
resultPackageVersion(result)
```

Arguments

result

a summarised results object

Value

a summarised results object

settings

Get settings from an object.

Description

Get settings from an object.

Usage

settings(x)

Arguments

Χ

Object

Value

A table with the settings of the object.

70 settings.cohort_table

settings.cohort_table Get cohort settings from a cohort_table object.

Description

Get cohort settings from a cohort_table object.

Usage

```
## S3 method for class 'cohort_table'
settings(x)
```

Arguments

Χ

A cohort_table object.

Value

A table with the details of the cohort settings.

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
person <- tibble(</pre>
  person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
  race_concept_id = 0, ethnicity_concept_id = 0
observation_period <- tibble(</pre>
  observation_period_id = 1, person_id = 1,
  observation_period_start_date = as.Date("2000-01-01"),
  observation_period_end_date = as.Date("2023-12-31"),
  period_type_concept_id = 0
)
cohort <- tibble(</pre>
  cohort_definition_id = 1,
  subject_id = 1,
  cohort_start_date = as.Date("2010-01-01"),
  cohort_end_date = as.Date("2012-01-01")
)
cdm <- cdmFromTables(</pre>
  tables = list("person" = person, "observation_period" = observation_period),
  cdmName = "test",
  cohortTables = list("my_cohort" = cohort)
)
settings(cdm$my_cohort)
cdm$my_cohort <- cdm$my_cohort |>
```

```
newCohortTable(cohortSetRef = tibble(
   cohort_definition_id = 1, cohort_name = "new_name"
))
settings(cdm$my_cohort)
```

settings.summarised_result

Get settings from a summarised_result object.

Description

Get settings from a summarised_result object.

Usage

```
## S3 method for class 'summarised_result'
settings(x)
```

Arguments

Х

A summarised_result object.

Value

A table with the settings.

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
person <- tibble(</pre>
  person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
  race_concept_id = 0, ethnicity_concept_id = 0
)
observation_period <- tibble(</pre>
  observation_period_id = 1, person_id = 1,
  observation_period_start_date = as.Date("2000-01-01"),
  observation_period_end_date = as.Date("2023-12-31"),
  period_type_concept_id = 0
)
cohort <- tibble(</pre>
  cohort_definition_id = 1,
  subject_id = 1,
  cohort_start_date = as.Date("2010-01-01"),
  cohort_end_date = as.Date("2012-01-01")
cdm <- cdmFromTables(</pre>
```

72 settingsColumns

```
tables = list("person" = person, "observation_period" = observation_period),
  cdmName = "test",
  cohortTables = list("my_cohort" = cohort)
)
result <- summary(cdm$my_cohort)
settings(result)</pre>
```

settingsColumns

Identify settings columns of a <summarised_result>

Description

Identifies and returns the columns of the settings table obtained by using settings() in a <summarised_result> object.

Usage

```
settingsColumns(result, metadata = FALSE)
```

Arguments

result A <summarised_result>.

metadata Whether to include metadata columns in settings or not.

Value

Vector with names of the settings columns

```
{
  library(dplyr)
  library(omopgenerics)

x <- tibble(
   "result_id" = as.integer(c(1, 2)),
   "cdm_name" = c("cprd", "eunomia"),
   "group_name" = "cohort_name",
   "group_level" = "my_cohort",
   "strata_name" = "sex",
   "strata_level" = "male",
   "variable_name" = "Age group",
   "variable_level" = "10 to 50",
   "estimate_name" = "count",
   "estimate_type" = "numeric",
   "estimate_value" = "5",</pre>
```

sourceType 73

```
"additional_name" = "overall",
    "additional_level" = "overall"
) |>
    newSummarisedResult(settings = tibble(
        "result_id" = c(1, 2), "custom" = c("A", "B")
    ))

x

x |> settingsColumns()
}
```

sourceType

Get the source type of an object.

Description

Get the source type of an object.

Usage

```
sourceType(x)
```

Arguments

Х

Object to know the source type.

Value

A character vector that defines the type of cdm_source.

splitAdditional

Split additional_name and additional_level columns

Description

Pivots the input dataframe so the values of the column additional_name are transformed into columns that contain values from the additional_level column.

Usage

```
splitAdditional(result, keep = FALSE, fill = "overall")
```

74 splitAll

Arguments

result A dataframe with at least the columns additional_name and additional_level.

keep Whether to keep the original group_name and group_level columns.

fill Optionally, a character that specifies what value should be filled in with when

missing.

Value

A dataframe.

Examples

```
{
 library(dplyr)
 library(omopgenerics)
 x <- tibble(
    "result_id" = as.integer(c(1, 2)),
    "cdm_name" = c("cprd", "eunomia"),
    "group_name" = "cohort_name",
    "group_level" = "my_cohort",
    "strata_name" = "sex",
    "strata_level" = "male",
    "variable_name" = "Age group",
    "variable_level" = "10 to 50",
    "estimate_name" = "count",
    "estimate_type" = "numeric",
    "estimate_value" = "5",
    "additional_name" = "overall",
    "additional_level" = "overall"
    newSummarisedResult(settings = tibble(
      "result_id" = c(1, 2), "custom" = c("A", "B")
 x |> splitAdditional()
```

splitAll

Split all pairs name-level into columns.

Description

Pivots the input dataframe so any pair name-level columns are transformed into columns (name) that contain values from the corresponding level.

splitAll 75

Usage

```
splitAll(result, keep = FALSE, fill = "overall", exclude = "variable")
```

Arguments

result A data.frame.

keep Whether to keep the original name-level columns.

fill A character that specifies what value should be filled in when missing.

exclude Name of a column pair to exclude.

Value

A dataframe with group, strata and additional as columns.

```
{
  library(dplyr)
  library(omopgenerics)
  x <- tibble(</pre>
    "result_id" = as.integer(c(1, 2)),
    "cdm_name" = c("cprd", "eunomia"),
    "group_name" = "cohort_name",
    "group_level" = "my_cohort",
    "strata_name" = "sex",
    "strata_level" = "male",
    "variable_name" = "Age group",
    "variable_level" = "10 to 50",
    "estimate_name" = "count",
    "estimate_type" = "numeric",
    "estimate_value" = "5",
    "additional_name" = "overall",
    "additional_level" = "overall"
    newSummarisedResult(settings = tibble(
      "result_id" = c(1, 2), "custom" = c("A", "B")
    ))
 x |> splitAll()
```

76 splitGroup

splitGroup

Split group_name and group_level columns

Description

Pivots the input dataframe so the values of the column group_name are transformed into columns that contain values from the group_level column.

Usage

```
splitGroup(result, keep = FALSE, fill = "overall")
```

Arguments

result A dataframe with at least the columns group_name and group_level.

keep Whether to keep the original group_name and group_level columns.

fill Optionally, a character that specifies what value should be filled in with when

Optionary, a character that specifies what value should be filled in with when

missing.

Value

A dataframe.

```
{
 library(dplyr)
 library(omopgenerics)
 x <- tibble(
    "result_id" = as.integer(c(1, 2)),
    "cdm_name" = c("cprd", "eunomia"),
    "group_name" = "cohort_name",
    "group_level" = "my_cohort",
    "strata_name" = "sex",
    "strata_level" = "male",
    "variable_name" = "Age group",
    "variable_level" = "10 to 50",
    "estimate_name" = "count",
    "estimate_type" = "numeric",
    "estimate_value" = "5",
    "additional_name" = "overall",
    "additional_level" = "overall"
   newSummarisedResult(settings = tibble(
      "result_id" = c(1, 2), "custom" = c("A", "B")
   ))
 Χ
```

splitStrata 77

```
x |> splitGroup()
}
```

splitStrata

Split strata_name and strata_level columns

Description

Pivots the input dataframe so the values of the column strata_name are transformed into columns that contain values from the strata_level column.

Usage

```
splitStrata(result, keep = FALSE, fill = "overall")
```

Arguments

result A dataframe with at least the columns strata_name and strata_level.

keep Whether to keep the original group_name and group_level columns.

Optionally, a character that specifies what value should be filled in with when missing.

Value

A dataframe.

```
library(dplyr)
library(omopgenerics)
x <- tibble(</pre>
  "result_id" = as.integer(c(1, 2)),
  "cdm_name" = c("cprd", "eunomia"),
  "group_name" = "cohort_name",
  "group_level" = "my_cohort",
  "strata_name" = "sex",
  "strata_level" = "male",
  "variable_name" = "Age group",
  "variable_level" = "10 to 50",
  "estimate_name" = "count",
  "estimate_type" = "numeric",
  "estimate_value" = "5",
  "additional_name" = "overall",
  "additional_level" = "overall"
) |>
  newSummarisedResult(settings = tibble(
```

78 strataColumns

```
"result_id" = c(1, 2), "custom" = c("A", "B")
))

x

x |> splitStrata()
}
```

strataColumns

Identify variables in strata_name column

Description

Identifies and returns the unique values in strata_name column.

Usage

```
strataColumns(result)
```

Arguments

result

A tibble.

Value

Unique values of the strata name column.

```
library(dplyr)
library(omopgenerics)
x <- tibble(</pre>
  "result_id" = as.integer(c(1, 2)),
  "cdm_name" = c("cprd", "eunomia"),
  "group_name" = "cohort_name",
  "group_level" = "my_cohort",
  "strata_name" = "sex",
  "strata_level" = "male",
  "variable_name" = "Age group",
  "variable_level" = "10 to 50",
  "estimate_name" = "count",
  "estimate_type" = "numeric",
  "estimate_value" = "5",
  "additional_name" = "overall",
  "additional_level" = "overall"
) |>
 newSummarisedResult(settings = tibble(
```

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```
"result_id" = c(1, 2), "custom" = c("A", "B")
))

x

x |> strataColumns()
}
```

summary.cdm_reference Summary a cdm reference

Description

Summary a cdm reference

Usage

```
## S3 method for class 'cdm_reference'
summary(object, ...)
```

Arguments

object A cdm reference object.
... For compatibility (not used).

Value

A summarised_result object with a summary of the data contained in the cdm.

```
library(dplyr, warn.conflicts = FALSE)
person <- tibble(</pre>
  person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
  race_concept_id = 0, ethnicity_concept_id = 0
)
observation_period <- tibble(</pre>
  observation_period_id = 1, person_id = 1,
  observation_period_start_date = as.Date("2000-01-01"),
  observation_period_end_date = as.Date("2023-12-31"),
  period_type_concept_id = 0
)
cdm <- cdmFromTables(</pre>
  tables = list("person" = person, "observation_period" = observation_period),
  cdmName = "test"
)
summary(cdm)
```

80 summary.cohort_table

```
summary.cohort_table Summary a generated cohort set
```

Description

Summary a generated cohort set

Usage

```
## S3 method for class 'cohort_table'
summary(object, ...)
```

Arguments

```
object A generated cohort set object.
... For compatibility (not used).
```

Value

A summarised_result object with a summary of a cohort_table.

```
library(dplyr, warn.conflicts = FALSE)
person <- tibble(</pre>
  person_id = 1, gender_concept_id = 0, year_of_birth = 1990,
  race_concept_id = 0, ethnicity_concept_id = 0
)
observation_period <- tibble(</pre>
  observation_period_id = 1, person_id = 1,
  observation_period_start_date = as.Date("2000-01-01"),
  observation_period_end_date = as.Date("2023-12-31"),
  period_type_concept_id = 0
)
cdm <- cdmFromTables(</pre>
  tables = list("person" = person, "observation_period" = observation_period),
  cdmName = "test",
  cohortTables = list("cohort1" = tibble(
    cohort_definition_id = 1,
    subject_id = 1,
    cohort_start_date = as.Date("2010-01-01"),
    cohort_end_date = as.Date("2010-01-05")
  ))
)
summary(cdm$cohort1)
```

```
\verb|summary.summarised_result|\\
```

Summary a summarised_result

Description

Summary a summarised_result

Usage

```
## S3 method for class 'summarised_result'
summary(object, ...)
```

Arguments

```
object A summarised_result object.
... For compatibility (not used).
```

Value

A summary of the result_types contained in a summarised_result object.

```
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("2023-12-31"),
    period_type_concept_id = 0
)

cdm <- cdmFromTables(
    tables = list("person" = person, "observation_period" = observation_period),
    cdmName = "test"
)

result <- summary(cdm)

summary(result)</pre>
```

suppress

Function to suppress counts in result objects

Description

Function to suppress counts in result objects

Usage

```
suppress(result, minCellCount = 5)
```

Arguments

result Result object

minCellCount Minimum count of records to report results.

Value

Table with suppressed counts

```
suppress.summarised_result
```

Function to suppress counts in result objects

Description

Function to suppress counts in result objects

Usage

```
## S3 method for class 'summarised_result'
suppress(result, minCellCount = 5)
```

Arguments

result summarised_result object.

minCellCount Minimum count of records to report results.

Value

summarised_result with suppressed counts.

tableName 83

Examples

```
library(dplyr, warn.conflicts = FALSE)
library(omopgenerics)
my_result <- tibble(</pre>
  "result_id" = "1",
  "cdm_name" = "mock",
  "result_type" = "summarised_characteristics",
  "package_name" = "omopgenerics",
  "package_version" = as.character(utils::packageVersion("omopgenerics")),
  "group_name" = "overall",
  "group_level" = "overall",
  "strata_name" = c(rep("overall", 6), rep("sex", 3)),
  "strata_level" = c(rep("overall", 6), "male", "female", "female"),
  "variable_name" = c(
    "number records", "age_group", "age_group",
    "age_group", "age_group", "my_variable", "number records", "age_group",
    "age_group"
  ),
  "variable_level" = c(
   NA, "<50", "<50", ">=50", ">=50", NA, NA,
    "<50", "<50"
  ),
  "estimate_name" = c(
    "count", "count", "percentage", "count", "percentage",
    "random", "count", "count", "percentage"
  ),
  "estimate_type" = c(
    "integer", "integer", "percentage", "integer",
    "percentage", "numeric", "integer", "integer", "percentage"
  "estimate_value" = c("10", "5", "50", "3", "30", "1", "3", "12", "6"),
  "additional_name" = "overall",
  "additional_level" = "overall"
)
my_result <- newSummarisedResult(my_result)</pre>
my_result |> glimpse()
my_result <- suppress(my_result, minCellCount = 5)</pre>
my_result |> glimpse()
```

tableName

Get the table name of a cdm_table.

Description

Get the table name of a cdm_table.

Usage

```
tableName(table)
```

84 tableSource

Arguments

table

A cdm_table.

Value

A character with the name.

Examples

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
cdm <- cdmFromTables(</pre>
  tables = list(
    "person" = tibble(
      person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
      race_concept_id = 0, ethnicity_concept_id = 0
   ),
    "observation_period" = tibble(
      observation_period_id = 1:3, person_id = 1:3,
      observation_period_start_date = as.Date("2000-01-01"),
      observation_period_end_date = as.Date("2023-12-31"),
      period_type_concept_id = 0
   )
  ),
  cdmName = "mock"
tableName(cdm$person)
```

tableSource

Get the table source of a cdm_table.

Description

Get the table source of a cdm_table.

Usage

```
tableSource(table)
```

Arguments

table

A cdm_table.

Value

A cdm_source object.

Examples

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
cdm <- cdmFromTables(</pre>
  tables = list(
    "person" = tibble(
      person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
      race_concept_id = 0, ethnicity_concept_id = 0
    "observation_period" = tibble(
      observation_period_id = 1:3, person_id = 1:3,
      observation_period_start_date = as.Date("2000-01-01"),
      observation_period_end_date = as.Date("2023-12-31"),
      period_type_concept_id = 0
   )
  ),
  cdmName = "mock"
tableSource(cdm$person)
```

tidy.summarised_result

Turn a <summarised_result> *object into a tidy tibble*

Description

[Experimental] Provides tools for obtaining a tidy version of a <summarised_result> object. This tidy version will include the settings as columns, estimate_value will be pivotted into columns using estimate_name as names, and group, strata, and additional will be splitted.

Usage

```
## S3 method for class 'summarised_result' tidy(x, ...)
```

Arguments

x A <summarised_result>.

... For compatibility (not used).

Value

A tibble.

86 tidyColumns

Examples

```
library(dplyr)
library(omopgenerics)
x <- tibble(</pre>
  "result_id" = as.integer(c(1, 2)),
  "cdm_name" = c("cprd", "eunomia"),
  "group_name" = "cohort_name",
  "group_level" = "my_cohort",
  "strata_name" = "sex",
  "strata_level" = "male",
  "variable_name" = "Age group",
  "variable_level" = "10 to 50",
  "estimate_name" = "count",
  "estimate_type" = "numeric",
  "estimate_value" = "5",
  "additional_name" = "overall",
  "additional_level" = "overall"
) |>
  newSummarisedResult(settings = tibble(
    "result_id" = c(1, 2), "custom" = c("A", "B")
  ))
Х
x \mid > tidy()
```

tidyColumns

Identify tidy columns of a <summarised_result>

Description

Identifies and returns the columns that the tidy version of the <summarised_result> will have.

Usage

```
tidyColumns(result)
```

Arguments

 $result \hspace{1cm} A < summarised_result >.$

Value

Table columns after applying tidy() function to a <summarised_result>.

tmpPrefix 87

Examples

```
library(dplyr)
library(omopgenerics)
x <- tibble(</pre>
  "result_id" = as.integer(c(1, 2)),
  "cdm_name" = c("cprd", "eunomia"),
  "group_name" = "cohort_name",
  "group_level" = "my_cohort",
  "strata_name" = "sex",
  "strata_level" = "male",
  "variable_name" = "Age group",
  "variable_level" = "10 to 50",
  "estimate_name" = "count",
  "estimate_type" = "numeric",
  "estimate_value" = "5",
  "additional_name" = "overall",
  "additional_level" = "overall"
) |>
  newSummarisedResult(settings = tibble(
    "result_id" = c(1, 2), "custom" = c("A", "B")
 ))
Х
x |> tidyColumns()
```

tmpPrefix

Create a temporary prefix for tables, that contains a unique prefix that starts with tmp.

Description

Create a temporary prefix for tables, that contains a unique prefix that starts with tmp.

Usage

```
tmpPrefix()
```

Value

A temporary prefix.

```
library(omopgenerics)
tmpPrefix()
```

88 uniqueId

toSnakeCase

Convert a character vector to snake case

Description

Convert a character vector to snake case

Usage

```
toSnakeCase(x)
```

Arguments

Х

Character vector to convert

Value

A snake_case vector

Examples

```
toSnakeCase("myVariable")
toSnakeCase(c("cohort1", "Cohort22b"))
```

uniqueId

Get a unique Identifier with a certain number of characters and a prefix.

Description

Get a unique Identifier with a certain number of characters and a prefix.

Usage

```
uniqueId(n = 1, exclude = character(), nChar = 3, prefix = "id_")
```

Arguments

n Number of identifiers.
 exclude Columns to exclude.
 nChar Number of characters.
 prefix A prefix for the identifiers.

Value

A character vector with n unique identifiers.

uniqueTableName 89

uniqueTableName

Create a unique table name

Description

Create a unique table name

Usage

```
uniqueTableName(prefix = "")
```

Arguments

prefix

Prefix for the table names.

Value

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

Examples

```
library(omopgenerics)
uniqueTableName()
```

uniteAdditional

Unite one or more columns in additional_name-additional_level format

Description

Unites targeted table columns into additional_name-additional_level columns.

Usage

```
uniteAdditional(
    x,
    cols = character(0),
    keep = FALSE,
    ignore = c(NA, "overall")
)
```

Arguments

x Tibble or dataframe.cols Columns to aggregate.

keep Whether to keep the original columns.

ignore Level values to ignore.

90 uniteGroup

Value

A tibble with the new columns.

Examples

```
x <- dplyr::tibble(
  variable = "number subjects",
  value = c(10, 15, 40, 78),
  sex = c("Male", "Female", "Male", "Female"),
  age_group = c("<40", ">40", ">40", "<40")
)

x |>
  uniteAdditional(c("sex", "age_group"))
```

uniteGroup

Unite one or more columns in group_name-group_level format

Description

Unites targeted table columns into group_name-group_level columns.

Usage

```
uniteGroup(x, cols = character(0), keep = FALSE, ignore = c(NA, "overall"))
```

Arguments

x Tibble or dataframe. cols Columns to aggregate.

keep Whether to keep the original columns.

ignore Level values to ignore.

Value

A tibble with the new columns.

```
x <- dplyr::tibble(
  variable = "number subjects",
  value = c(10, 15, 40, 78),
  sex = c("Male", "Female", "Male", "Female"),
  age_group = c("<40", ">40", ">40", "<40")
)
x |>
  uniteGroup(c("sex", "age_group"))
```

uniteStrata 91

uniteStrata

Unite one or more columns in strata_name-strata_level format

Description

Unites targeted table columns into strata_name-strata_level columns.

Usage

```
uniteStrata(x, cols = character(0), keep = FALSE, ignore = c(NA, "overall"))
```

Arguments

x Tibble or dataframe.cols Columns to aggregate.

keep Whether to keep the original columns.

ignore Level values to ignore.

Value

A tibble with the new columns.

Examples

```
x <- dplyr::tibble(
  variable = "number subjects",
  value = c(10, 15, 40, 78),
  sex = c("Male", "Female", "Male", "Female"),
  age_group = c("<40", ">40", ">40", "<40")
)
x |>
  uniteStrata(c("sex", "age_group"))
```

validate Age Group Argument

validateAgeGroupArgument

Description

validateAgeGroupArgument

Usage

```
validateAgeGroupArgument(
  ageGroup,
  multipleAgeGroup = TRUE,
  overlap = FALSE,
  null = TRUE,
  ageGroupName = "age_group",
  call = parent.frame()
)
```

Arguments

Value

validate ageGroup

validateCdmArgument

Description

validate Cdm Argument

Usage

```
validateCdmArgument(
  cdm,
  checkOverlapObservation = FALSE,
  checkStartBeforeEndObservation = FALSE,
  checkPlausibleObservationDates = FALSE,
  checkPerson = FALSE,
  validation = "error",
  call = parent.frame()
)
```

validateCohortArgument

Arguments

cdm A cdm_reference object

checkOverlapObservation

TRUE to perform check on no overlap observation period

checkStartBeforeEndObservation

TRUE to perform check on correct observational start and end date

checkPlausibleObservationDates

TRUE to perform check that there are no implausible observation period start

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dates (before 1800-01-01) or end dates (after the current date)

checkPerson TRUE to perform check on person id in all clincial table are in person table

validation How to perform validation: "error", "warning".

call A call argument to pass to cli functions.

Value

A cdm_reference object

validateCohortArgument

Validate a cohort table input.

Description

Validate a cohort table input.

Usage

```
validateCohortArgument(
  cohort,
  checkEndAfterStart = FALSE,
  checkOverlappingEntries = FALSE,
  checkMissingValues = FALSE,
  checkInObservation = FALSE,
  dropExtraColumns = FALSE,
  validation = "error",
  call = parent.frame()
)
```

Arguments

cohort Object to be validated as a valid cohort input.

checkEndAfterStart

If TRUE a check that all cohort end dates come on or after cohort start date will be performed.

checkOverlappingEntries

If TRUE a check that no individuals have overlapping cohort entries will be performed.

checkMissingValues

If TRUE a check that there are no missing values in required fields will be performed.

checkInObservation

If TRUE a check that cohort entries are within the individuals observation periods will be performed.

dropExtraColumns

Whether to drop extra columns that are not the required ones.

validation How to perform validation: "error", "warning".

call A call argument to pass to cli functions.

validateCohortIdArgument

Validate cohortId argument. CohortId can either be a cohort_definition_id value, a cohort_name or a tidyselect expression referinc to cohort_names. If you want to support tidyselect expressions please use the function as: validateCohortIdArgument({{cohortId}}, cohort).

Description

Validate cohortId argument. CohortId can either be a cohort_definition_id value, a cohort_name or a tidyselect expression referinc to cohort_names. If you want to support tidyselect expressions please use the function as: validateCohortIdArgument({{cohortId}}, cohort).

Usage

```
validateCohortIdArgument(
  cohortId,
  cohort,
  validation = "error",
  call = parent.frame()
)
```

Arguments

cohortId A cohortId vector to be validated.

cohort A cohort_table object.

validation How to perform validation: "error", "warning".

call A call argument to pass to cli functions.

 $validate {\tt ConceptSetArgument}$

Validate conceptSet argument.

Description

Validate conceptSet argument.

Usage

```
validateConceptSetArgument(
  conceptSet,
  cdm = NULL,
  validation = "error",
  call = parent.frame()
)
```

Arguments

conceptSet It can be either a named list of concepts or a codelist, codelist_with_details or

conceptSetExpression object.

cdm A cdm_reference object, needed if a conceptSetExpression is provided.

validation How to perform validation: "error", "warning".

call A call argument to pass to cli functions.

validateNameArgument Validate name argument.

Description

Validate name argument.

Usage

```
validateNameArgument(
  name,
  cdm = NULL,
  validation = "error",
  null = FALSE,
  call = parent.frame()
)
```

96 validateNameLevel

Arguments

name Name of a new table to be added to a cdm object.

cdm A cdm_reference object. It will check if a table named name already exists in

the cdm.

validation How to perform validation: "error", "warning".

null If TRUE, name can be NULL

call A call argument to pass to cli functions.

validateNameLevel

Validate if two columns are valid Name-Level pair.

Description

Validate if two columns are valid Name-Level pair.

Usage

```
validateNameLevel(
   x,
   prefix,
   sep = " &&& ",
   validation = "error",
   call = parent.frame()
)
```

Arguments

x A tibble.

prefix Prefix for the name-level pair, e.g. 'strata' for strata_name-strata_level pair.

sep Separation pattern.

validation Either 'error', 'warning' or 'message'.

call Will be used by cli to report errors.

validateNewColumn 97

validateNewColumn	Validate a new column of a table
Validatenteweolumi	variable a new committee of a table

Description

Validate a new column of a table

Usage

```
validateNewColumn(table, column, validation = "warning", call = parent.frame())
```

Arguments

table The table to check if the column already exists.

column Character vector with the name(s) of the new column(s).

validation Whether to throw warning or error.

call Passed to cli functions.

Value

table without conflicting columns.

```
{\tt validateResultArguemnt}
```

validateResultArgument

Description

validate Result Argument

Usage

```
validateResultArguemnt(result, validation = "error", call = parent.frame())
```

Arguments

result summarise result object to validate

validation message to return call parent.frame

Value

summarise result object

```
validateResultArgument
```

validateResultArgument

Description

validateResultArgument

Usage

```
validateResultArgument(
  result,
  checkNoDuplicates = FALSE,
  checkSuppression = FALSE,
  validation = "error",
  call = parent.frame()
)
```

Arguments

result summarised_result object to validate.

checkNoDuplicates

Whether there are not allowed duplicates in the result object.

checkNameLevel Whether the name-level paired columns are can be correctly split. checkSuppression

Whether the suppression in the result object is well defined.

validation Only error is supported at the moment.

call parent.frame

Value

summarise result object

validateWindowArgument

validateWindowArgument

Description

validateWindowArgument

Usage

```
validateWindowArgument(window, snakeCase = TRUE, call = parent.frame())
```

[[.cdm_reference 99

Arguments

window time window

snakeCase return default window name in snake case if TRUE

call A call argument to pass to cli functions.

Value

time window

[[.cdm_reference

Subset a cdm reference object.

Description

Subset a cdm reference object.

Usage

```
## S3 method for class 'cdm_reference'
x[[name]]
```

Arguments

x A cdm reference

name The name or index of the table to extract from the cdm object.

Value

A single cdm table reference

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

cdm <- cdmFromTables(
   tables = list(
    "person" = tibble(
        person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
        race_concept_id = 0, ethnicity_concept_id = 0
   ),
    "observation_period" = tibble(
        observation_period_id = 1:3, person_id = 1:3,
        observation_period_start_date = as.Date("2000-01-01"),
        observation_period_end_date = as.Date("2023-12-31"),
        period_type_concept_id = 0
   )
   ),</pre>
```

\$.cdm_reference

```
cdmName = "mock"
)
cdm[["person"]]
```

[[<-.cdm_reference

Assign a table to a cdm reference.

Description

Assign a table to a cdm reference.

Usage

```
## S3 replacement method for class 'cdm_reference'
cdm[[name]] <- value</pre>
```

Arguments

cdm A cdm reference.

name Name where to assign the new table.

value Table with the same source than the cdm object.

Value

The cdm reference.

\$.cdm_reference

Subset a cdm reference object.

Description

Subset a cdm reference object.

Usage

```
## S3 method for class 'cdm_reference'
x$name
```

Arguments

x A cdm reference.

name The name of the table to extract from the cdm object.

\$<-.cdm_reference

Value

A single cdm table reference

Examples

```
library(omopgenerics)
library(dplyr, warn.conflicts = FALSE)
cdm <- cdmFromTables(</pre>
 tables = list(
    "person" = tibble(
      person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
      race_concept_id = 0, ethnicity_concept_id = 0
   ),
    "observation_period" = tibble(
      observation_period_id = 1:3, person_id = 1:3,
      observation_period_start_date = as.Date("2000-01-01"),
      observation_period_end_date = as.Date("2023-12-31"),
      period_type_concept_id = 0
   )
 ),
 cdmName = "mock"
)
cdm$person
```

\$<-.cdm_reference</pre>

Assign an table to a cdm reference.

Description

Assign an table to a cdm reference.

Usage

```
## S3 replacement method for class 'cdm_reference'
cdm$name <- value</pre>
```

Arguments

cdm A cdm reference.

name Name where to assign the new table.

value Table with the same source than the cdm object.

Value

The cdm reference.

\$<-.cdm_reference

```
library(omopgenerics)
cdm <- cdmFromTables(</pre>
  tables = list(
    "person" = dplyr::tibble(
     person_id = c(1, 2, 3), gender_concept_id = 0, year_of_birth = 1990,
     race_concept_id = 0, ethnicity_concept_id = 0
    "observation_period" = dplyr::tibble(
      observation_period_id = 1:3, person_id = 1:3,
      observation_period_start_date = as.Date("2000-01-01"),
      observation_period_end_date = as.Date("2023-12-31"),
      period_type_concept_id = 0
   )
 ),
  cdmName = "mock"
)
cdm$person
```

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