## Package 'CodelistGenerator'

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Title Identify Relevant Clinical Codes and Evaluate Their Use

Version 3.2.1 **Description** Generate a candidate code list for the Observational Medical Outcomes Partnership (OMOP) common data model based on string matching. For a given search strategy, a candidate code list will be returned. **License** Apache License (>= 2) **Encoding UTF-8** RoxygenNote 7.3.2 **Depends** R (>= 3.5.0) **Imports** checkmate (>= 2.0.0), DBI (>= 1.1.0), dplyr (>= 1.1.0), magrittr (>= 2.0.0), omopgenerics (>= 0.2.2), rlang (>= 1.0.0), glue (>= 1.5.0), stringr (>= 1.4.0), tidyselect (>= 1.2.0), tidyr (>= 1.2.0), cli (>= 3.1.0), purrr, lubridate, PatientProfiles (>= 1.1.0), RJSONIO, vctrs, visOmopResults (>= 0.3.0), lifecycle Suggests covr, duckdb, CDMConnector (>= 1.3.0), knitr, rmarkdown, testthat (>= 3.0.0), RPostgres, odbc, spelling, tibble, gt, flextable Config/testthat/edition 3 Config/testthat/parallel true VignetteBuilder knitr URL https://darwin-eu.github.io/CodelistGenerator/ Language en-US LazyData true **NeedsCompilation** no **Author** Edward Burn [aut, cre] (<a href="https://orcid.org/0000-0002-9286-1128">https://orcid.org/0000-0002-9286-1128</a>), Marti Catala [ctb] (<https://orcid.org/0000-0003-3308-9905>), Xihang Chen [aut] (<a href="https://orcid.org/0009-0001-8112-8959">https://orcid.org/0009-0001-8112-8959</a>), Nuria Mercade-Besora [aut] (<a href="https://orcid.org/0009-0006-7948-3747">https://orcid.org/0009-0006-7948-3747</a>), Mike Du [ctb] (<https://orcid.org/0000-0002-9517-8834>), Danielle Newby [ctb] (<a href="https://orcid.org/0000-0002-3001-1478">https://orcid.org/0000-0002-3001-1478</a>)

2 Contents

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## **Contents**

Index

availableATC
availableICD10
availableIngredients
codesFromCohort
codesFromConceptSet
codesInUse
compareCodelists
doseFormToRoute
getATCCodes
getCandidateCodes
getConceptClassId
getDescendants
getDomains
getDoseForm
getDoseUnit
getDrugIngredientCodes
getICD10StandardCodes
getMappings
getRelationshipId
getRouteCategories
getVocabularies
getVocabVersion
mockVocabRef
restrictToCodesInUse
sourceCodesInUse
stratifyByConcept
stratifyByDoseUnit
stratifyByRouteCategory
subsetOnDoseUnit
subsetOnRouteCategory
subsetToCodesInUse
summariseAchillesCodeUse
summariseCodeUse
summariseCohortCodeUse
summariseOrphanCodes
tableAchillesCodeUse
tableCodeUse
tableCohortCodeUse
tableOrphanCodes
=

**35** 

availableATC 3

availableATC

Get all ATC codes from the cdm

#### **Description**

Get all ATC codes from the cdm

#### Usage

```
availableATC(cdm, level = c("ATC 1st"))
```

## **Arguments**

cdm\_reference via CDMConnector

level ATC level. Can be one or more of "ATC 1st", "ATC 2nd", "ATC 3rd", "ATC

4th", and "ATC 5th"

#### Value

A vector list of all ATC codes for the chosen level(s) found in the concept table of cdm.

## **Examples**

```
cdm <- mockVocabRef()
availableATC(cdm)</pre>
```

availableICD10

Get all ICD codes from the cdm

#### **Description**

Get all ICD codes from the cdm

## Usage

```
availableICD10(cdm, level = c("ICD10 Chapter", "ICD10 SubChapter"))
```

## **Arguments**

cdm\_reference via CDMConnector

level Can be either "ICD10 Chapter" or "ICD10 SubChapter"

#### Value

A vector list of all ICD10 codes for the chosen level(s) found in the concept table of cdm.

4 codesFromCohort

#### **Examples**

```
cdm <- mockVocabRef()
availableICD10(cdm)</pre>
```

availableIngredients Get all ingredients codes from the cdm

## Description

Get all ingredients codes from the cdm

#### Usage

```
availableIngredients(cdm)
```

#### **Arguments**

cdm

cdm\_reference via CDMConnector

#### Value

A vector list of all ingredient level codes found in the concept table of cdm.

#### **Examples**

```
cdm <- mockVocabRef()
availableIngredients(cdm)</pre>
```

codesFromCohort

Get concept ids from a provided path to cohort json files

#### Description

Get concept ids from a provided path to cohort json files

#### Usage

```
codesFromCohort(path, cdm, type = c("codelist"))
```

#### **Arguments**

path	Path to a file or	r folder containing	JSONs of cohort	definitions

cdm A cdm reference created with CDMConnector

type Can be "codelist", "codelist\_with\_details", or "concept\_set\_expression"

codesFromConceptSet 5

#### Value

Named list with concept\_ids for each concept set

codesFromConceptSet

Get concept ids from a provided path to json files

## Description

Get concept ids from a provided path to json files

## Usage

```
codesFromConceptSet(path, cdm, type = c("codelist"))
```

#### **Arguments**

path Path to a file or folder containing JSONs of concept sets

cdm A cdm reference created with CDMConnector

type Can be "codelist", "codelist\_with\_details", or "concept\_set\_expression"

#### Value

Named list with concept\_ids for each concept set

#### **Examples**

codesInUse

Use achilles counts to get codes used in the database

## Description

Use achilles counts to get codes used in the database

6 compareCodelists

#### Usage

```
codesInUse(
  cdm,
  minimumCount = 0,
  table = c("condition_occurrence", "device_exposure", "drug_exposure", "measurement",
        "observation", "procedure_occurrence", "visit_occurrence")
)
```

#### **Arguments**

cdm\_reference via CDMConnector

minimumCount Any codes with a frequency under this will be removed.

table cdm table

#### Value

A list of integers indicating codes being used in the database.

## **Examples**

```
cdm <- mockVocabRef("database")
x <- codesInUse(cdm = cdm)
x
CDMConnector::cdmDisconnect(cdm)</pre>
```

compareCodelists

Compare two codelists

#### **Description**

Compare two codelists

## Usage

```
compareCodelists(codelist1, codelist2)
```

## **Arguments**

codelist1 Output of getCandidateCodes codelist2 Output of getCandidateCodes

#### Value

tibble

doseFormToRoute 7

#### **Examples**

```
cdm <- mockVocabRef()
codes1 <- getCandidateCodes(
  cdm = cdm,
  keywords = "Arthritis",
  domains = "Condition",
  includeDescendants = TRUE
)

codes2 <- getCandidateCodes(
  cdm = cdm,
  keywords = c("knee osteoarthritis", "arthrosis"),
  domains = "Condition",
  includeDescendants = TRUE
)

compareCodelists(
  codelist1 = codes1,
  codelist2 = codes2
)</pre>
```

 ${\tt doseFormToRoute}$ 

Equivalence from dose from concept IDs to route categories.

#### **Description**

Equivalence from dose from concept IDs to route categories.

## Usage

doseFormToRoute

#### **Format**

A data frame with two variables: dose\_form\_concept\_id and route\_category.

getATCCodes

Get descendant codes for ATC levels

#### **Description**

Get descendant codes for ATC levels

8 getCandidateCodes

#### Usage

```
getATCCodes(
  cdm,
  level = c("ATC 1st"),
  name = NULL,
  doseForm = NULL,
  doseUnit = NULL,
  routeCategory = NULL,
  type = "codelist"
)
```

# **Arguments** cdm

level	ATC level. Can be one or more of "ATC 1st", "ATC 2nd", "ATC 3rd", "ATC 4th", and "ATC 5th" $$
name	ATC name of interest. For example, c("Dermatologicals", "Nervous System"), would result in a list of length two with the descendant concepts for these two particular ATC groups.
doseForm	Only descendants codes with the specified dose form will be returned. If NULL, descendant codes will be returned regardless of dose form.
doseUnit	Only descendants codes with the specified dose unit will be returned. If NULL,

cdm\_reference via CDMConnector

Only descendants codes with the specified dose unit will be returned. If NULL,

descendant codes will be returned regardless of dose unit

routeCategory Only descendants codes with the specified route will be returned. If NULL,

descendant codes will be returned regardless of dose form.

type Can be "codelist", "codelist\_with\_details", or "concept\_set\_expression"

#### Value

Concepts with their format based on the type argument.

#### **Examples**

```
cdm <- mockVocabRef()
getATCCodes(cdm = cdm, level = "ATC 1st")</pre>
```

 ${\tt getCandidateCodes}$ 

Generate candidate codelist for the OMOP CDM

#### **Description**

This function generates a set of codes that can be considered for creating a phenotype using the OMOP CDM.

getCandidateCodes 9

#### Usage

```
getCandidateCodes(
  cdm,
  keywords,
  exclude = NULL,
  domains = "Condition",
  standardConcept = "Standard",
  searchInSynonyms = FALSE,
  searchNonStandard = FALSE,
  includeDescendants = TRUE,
  includeAncestor = FALSE
)
```

#### **Arguments**

cdm\_reference via CDMConnector

keywords Character vector of words to search for. Where more than one word is given

(e.g. "knee osteoarthritis"), all combinations of those words should be identified

positions (e.g. "osteoarthritis of knee") should be identified.

exclude Character vector of words to identify concepts to exclude.

domains Character vector with one or more of the OMOP CDM domain.

standardConcept

Character vector with one or more of "Standard", "Classification", and "Non-standard". These correspond to the flags used for the standard\_concept field in the concept table of the cdm.

searchInSynonyms

Either TRUE or FALSE. If TRUE the code will also search using both the primary name in the concept table and synonyms from the concept synonym table.

searchNonStandard

Either TRUE or FALSE. If TRUE the code will also search via non-standard concepts.

includeDescendants

Either TRUE or FALSE. If TRUE descendant concepts of identified concepts will be included in the candidate codelist.

includeAncestor

Either TRUE or FALSE. If TRUE the direct ancestor concepts of identified concepts will be included in the candidate codelist.

#### Value

tibble

#### **Examples**

```
cdm <- CodelistGenerator::mockVocabRef()
CodelistGenerator::getCandidateCodes(
   cdm = cdm,</pre>
```

10 getDescendants

```
keywords = "osteoarthritis"
)
```

 ${\tt getConceptClassId}$ 

getConceptClassId

## Description

getConceptClassId

#### Usage

```
getConceptClassId(cdm, standardConcept = "Standard", domain = NULL)
```

## Arguments

cdm\_reference via CDMConnector

standardConcept

Character vector with one or more of "Standard", "Classification", and "Non-standard". These correspond to the flags used for the standard\_concept field in

the concept table of the cdm.

domain Vocabulary domain

#### Value

The concept class used for a given set of domains

## **Examples**

```
cdm <- mockVocabRef()
getConceptClassId(cdm = cdm, domain = "drug")</pre>
```

getDescendants

getDescendants

#### **Description**

getDescendants

getDomains 11

#### Usage

```
getDescendants(
  cdm,
  conceptId,
  withAncestor = FALSE,
  ingredientRange = c(0, Inf),
  doseForm = NULL
)
```

#### **Arguments**

cdm\_reference via CDMConnector

conceptId concpet\_id to search

withAncestor If TRUE, return column with ancestor. In case of multiple ancestors, concepts

will be separated by ";"

ingredientRange

Used to restrict descendant codes to those associated with a specific number of drug ingredients. Must be a vector of length two with the first element the minimum number of ingredients allowed and the second the maximum. A value of c(2, 2) would restrict to only concepts associated with two ingredients.

doseForm

Only descendants codes with the specified drug dose form will be returned. If

NULL, descendant codes will be returned regardless of dose form.

#### Value

The descendants of a given concept id

#### **Examples**

```
cdm <- mockVocabRef()
getDescendants(cdm = cdm, conceptId = 1)</pre>
```

getDomains

getDomains

## Description

getDomains

#### Usage

```
getDomains(cdm, standardConcept = "Standard")
```

12 getDoseForm

#### **Arguments**

cdm\_reference via CDMConnector

standardConcept

Character vector with one or more of "Standard", "Classification", and "Non-standard". These correspond to the flags used for the standard\_concept field in the concept table of the cdm.

#### Value

The domains of the cdm

## Examples

```
cdm <- mockVocabRef()
getDomains(cdm = cdm)</pre>
```

getDoseForm

getDoseForm

## Description

getDoseForm

#### Usage

```
getDoseForm(cdm)
```

## **Arguments**

cdm

cdm\_reference via CDMConnector

## Value

The dose forms available for drug concepts

## **Examples**

```
cdm <- mockVocabRef()
getDoseForm(cdm = cdm)</pre>
```

getDoseUnit 13

getDoseUnit

Get available routes in a cdm reference.

#### **Description**

Get the dose form categories available in the database (see https://doi.org/10.1002/pds.5809) for more details on how routes were classified).

## Usage

```
getDoseUnit(cdm)
```

#### **Arguments**

cdm

A cdm reference.

#### Value

A character vector with available routes

```
getDrugIngredientCodes
```

Get descendant codes for drug ingredients

## Description

Get descendant codes for drug ingredients

## Usage

```
getDrugIngredientCodes(
   cdm,
   name = NULL,
   nameStyle = "{concept_code}_{concept_name}",
   doseForm = NULL,
   doseUnit = NULL,
   routeCategory = NULL,
   ingredientRange = c(1, Inf),
   type = "codelist"
)
```

#### **Arguments**

cdmcdm\_reference via CDMConnector name Names of ingredients of interest. For example, c("acetaminophen", "codeine"), would result in a list of length two with the descendant concepts for these two particular drug ingredients. nameStyle Name style to apply to returned list. Can be one of "{concept\_code}\_{concept\_name}", "{concept\_code}", or "{concept\_name}". doseForm Only descendants codes with the specified dose form will be returned. If NULL, descendant codes will be returned regardless of dose form. doseUnit Only descendants codes with the specified dose unit will be returned. If NULL, descendant codes will be returned regardless of dose unit Only descendants codes with the specified route will be returned. If NULL, routeCategory descendant codes will be returned regardless of route category. ingredientRange Used to restrict descendant codes to those associated with a specific number of ingredients. Must be a vector of length two with the first element the minimum number of ingredients allowed and the second the maximum. A value of c(2, 2)would restrict to only concepts associated with two ingredients. Can be "codelist", "codelist\_with\_details", or "concept\_set\_expression" type

#### Value

Concepts with their format based on the type argument.

## **Examples**

 ${\it getICD10StandardCodes} \begin{tabular}{ll} \it Get \ corresponding \ standard \ codes \ for \ \it ICD-10 \ chapters \ and \ sub-chapters \end{tabular}$ 

#### **Description**

Get corresponding standard codes for ICD-10 chapters and sub-chapters

getMappings 15

#### Usage

```
getICD10StandardCodes(
  cdm,
  level = c("ICD10 Chapter", "ICD10 SubChapter"),
  name = NULL,
  includeDescendants = TRUE,
  type = "codelist"
)
```

#### **Arguments**

cdm\_reference via CDMConnector

level Can be either "ICD10 Chapter" or "ICD10 SubChapter"

name Name of chapter or sub-chapter of interest. If NULL, all will be considered.

includeDescendants

If FALSE only direct mappings from ICD-10 codes to standard codes will be returned. If TRUE descendants of standard concepts will also be included.

type Can be "codelist", "codelist\_with\_details", or "concept\_set\_expression"

#### Value

A named list, with each element containing the corresponding standard codes (and descendants) of ICD chapters and sub-chapters

#### **Examples**

```
cdm <- mockVocabRef()
getICD10StandardCodes(cdm = cdm, level = c(
  "ICD10 Chapter",
   "ICD10 SubChapter"
))</pre>
```

getMappings

Show mappings from non-standard vocabularies to standard

#### **Description**

Show mappings from non-standard vocabularies to standard

## Usage

```
getMappings(
  candidateCodelist,
  cdm = NULL,
  nonStandardVocabularies = c("ATC", "ICD10CM", "ICD10PCS", "ICD9CM", "ICD9Proc",
      "LOINC", "OPCS4", "Read", "RxNorm", "RxNorm Extension", "SNOMED")
)
```

16 getRelationshipId

## **Arguments**

#### Value

tibble

## **Examples**

```
cdm <- CodelistGenerator::mockVocabRef()
codes <- CodelistGenerator::getCandidateCodes(
   cdm = cdm,
   keywords = "osteoarthritis"
)
CodelistGenerator::getMappings(
   cdm = cdm,
   candidateCodelist = codes,
   nonStandardVocabularies = "READ"
)</pre>
```

getRelationshipId

Get relationship ID values from the concept relationship table

## **Description**

Get relationship ID values from the concept relationship table

## Usage

```
getRelationshipId(
  cdm,
  standardConcept1 = "standard",
  standardConcept2 = "standard",
  domains1 = "condition",
  domains2 = "condition"
)
```

## **Arguments**

 $\mathsf{cdm}$ 

A cdm reference

getRouteCategories 17

standardConcept1

Character vector with one or more of "Standard", "Classification", and "Non-standard". These correspond to the flags used for the standard\_concept field in the concept table of the cdm.

standardConcept2

Character vector with one or more of "Standard", "Classification", and "Non-standard". These correspond to the flags used for the standard\_concept field in the concept table of the cdm.

domains1 Character vector with one or more of the OMOP CDM domain.

domains2 Character vector with one or more of the OMOP CDM domain.

#### Value

A character vector with unique values

### **Examples**

```
cdm <- mockVocabRef()
getRelationshipId(cdm = cdm)</pre>
```

getRouteCategories

Get available routes in a cdm reference.

## Description

Get the dose form categories available in the database (see https://doi.org/10.1002/pds.5809) for more details on how routes were classified).

#### Usage

```
getRouteCategories(cdm)
```

#### **Arguments**

cdm

A cdm reference.

#### Value

A character vector with available routes

18 getVocabVersion

getVocabularies

getVocabularies

## Description

getVocabularies

## Usage

```
getVocabularies(cdm)
```

## Arguments

cdm

cdm\_reference via CDMConnector

## Value

Names of available vocabularies

## **Examples**

```
cdm <- mockVocabRef()
getVocabularies(cdm = cdm)</pre>
```

getVocabVersion

getVocabVersion

## Description

getVocabVersion

#### Usage

```
getVocabVersion(cdm)
```

## Arguments

cdm

cdm\_reference via CDMConnector

## Value

the vocabulary version being used

## **Examples**

```
cdm <- mockVocabRef()
getVocabVersion(cdm = cdm)</pre>
```

mockVocabRef 19

mockVocabRef

Generate example vocabulary database

#### **Description**

Generate example vocabulary database

#### Usage

```
mockVocabRef(backend = "data_frame")
```

## Arguments

backend

'database' (duckdb) or 'data\_frame'

#### Value

cdm reference with mock vocabulary

## **Examples**

```
cdm <- mockVocabRef()</pre>
cdm
```

the database

## Description

Use achilles counts to filter a codelist to keep only the codes used in the database

#### Usage

```
restrictToCodesInUse(
  х,
 cdm,
 minimumCount = 0L,
 table = c("condition_occurrence", "device_exposure", "drug_exposure", "measurement",
    "observation", "procedure_occurrence", "visit_occurrence")
)
```

20 sourceCodesInUse

#### **Arguments**

x A codelist

cdm cdm\_reference via CDMConnector

minimumCount Any codes with a frequency under this will be removed.

table cdm table

#### Value

Use achilles counts to filter codelist to only the codes used in the database

#### **Examples**

sourceCodesInUse

Use achilles counts to get source codes used in the database

#### **Description**

Use achilles counts to get source codes used in the database

## Usage

#### **Arguments**

cdm\_reference via CDMConnector

table cdm table

#### Value

A list of source codes used in the database.

stratifyByConcept 21

#### **Examples**

```
cdm <- mockVocabRef("database")
x <- sourceCodesInUse(cdm = cdm)
x
CDMConnector::cdmDisconnect(cdm)</pre>
```

stratifyByConcept

Stratify a codelist by the concepts included within it

#### **Description**

Stratify a codelist by the concepts included within it

#### Usage

```
stratifyByConcept(x, cdm, keepOriginal = FALSE)
```

#### **Arguments**

x A codelist

cdm A cdm reference

keepOriginal Whether to keep the original codelist and append the stratify (if TRUE) or just

return the stratified codelist (if FALSE).

#### Value

A codelist

stratifyByDoseUnit

Stratify a codelist by dose unit

#### **Description**

Stratify a codelist by dose unit

## Usage

```
stratifyByDoseUnit(x, cdm, keepOriginal = FALSE)
```

## Arguments

x A codelist

cdm A cdm reference

keepOriginal Whether to keep the original codelist and append the stratify (if TRUE) or just

return the stratified codelist (if FALSE).

22 subsetOnDoseUnit

#### Value

A codelist

stratifyByRouteCategory

Stratify a codelist by route category

## Description

Stratify a codelist by route category

#### Usage

```
stratifyByRouteCategory(x, cdm, keepOriginal = FALSE)
```

## Arguments

x A codelist

cdm A cdm reference

keepOriginal Whether to keep the original codelist and append the stratify (if TRUE) or just

return the stratified codelist (if FALSE).

#### Value

A codelist

subsetOnDoseUnit Subset a codelist to only those with a particular dose unit

#### **Description**

Subset a codelist to only those with a particular dose unit

### Usage

```
subsetOnDoseUnit(x, cdm, doseUnit)
```

#### **Arguments**

x Codelist

cdm A cdm reference

doseUnit Dose unit. Use getDoseUnit() to find the available dose units in a cdm

#### Value

The codelist with only those concepts associated with the dose unit

subsetOnRouteCategory Subset a codelist to only those with a particular route category

#### **Description**

Subset a codelist to only those with a particular route category

## Usage

```
subsetOnRouteCategory(x, cdm, routeCategory)
```

#### **Arguments**

x Codelist

cdm A cdm reference

routeCategory Route category. Use getRoutes() to find the available route categories for a cdm

#### Value

The codelist with only those concepts associated with the specified route categories

## Description

Use achilles counts to filter a codelist to keep only the codes used in the database

## Usage

```
subsetToCodesInUse(
    x,
    cdm,
    minimumCount = 0L,
    table = c("condition_occurrence", "device_exposure", "drug_exposure", "measurement",
        "observation", "procedure_occurrence", "visit_occurrence")
)
```

#### **Arguments**

x A codelist

cdm\_reference via CDMConnector

minimumCount Any codes with a frequency under this will be removed.

table cdm table

#### Value

Use achilles counts to filter codelist to only the codes used in the database

## **Examples**

summariseAchillesCodeUse

Summarise code use from achilles counts

## Description

Summarise code use from achilles counts

#### Usage

```
summariseAchillesCodeUse(
    x,
    cdm,
    countBy = c("record", "person"),
    minCellCount = lifecycle::deprecated()
)
```

## **Arguments**

#### Value

A tibble with results

summariseCodeUse 25

#### **Examples**

```
cdm <- mockVocabRef("database")
oa <- getCandidateCodes(cdm = cdm, keywords = "osteoarthritis")
result_achilles <- summariseAchillesCodeUse(list(oa = oa$concept_id), cdm = cdm)
result_achilles
CDMConnector::cdmDisconnect(cdm)</pre>
```

summariseCodeUse

Summarise code use in patient-level data

#### **Description**

Summarise code use in patient-level data

## Usage

```
summariseCodeUse(
    x,
    cdm,
    countBy = c("record", "person"),
    byConcept = TRUE,
    byYear = FALSE,
    bySex = FALSE,
    ageGroup = NULL,
    minCellCount = lifecycle::deprecated()
)
```

## Arguments

x List of concept IDs

cdm\_reference via CDMConnector::cdm\_from\_con()

countBy Either "record" for record-level counts or "person" for person-level counts

byConcept TRUE or FALSE. If TRUE code use will be summarised by

by Year TRUE or FALSE. If TRUE code use will be summarised by year.

by Sex TRUE or FALSE. If TRUE code use will be summarised by sex.

ageGroup If not NULL, a list of ageGroup vectors of length two.

minCellCount \ifelse{html}{\href{https://lifecycle.r-lib.org/articles/stages.html#deprecated}{\fi

#### Value

A tibble with results overall and, if specified, by strata

## **Examples**

summariseCohortCodeUse

Summarise code use among a cohort in the cdm reference

#### Description

Summarise code use among a cohort in the cdm reference

#### Usage

```
summariseCohortCodeUse(
    x,
    cdm,
    cohortTable,
    cohortId = NULL,
    timing = "any",
    countBy = c("record", "person"),
    byConcept = TRUE,
    byYear = FALSE,
    bySex = FALSE,
    ageGroup = NULL,
    minCellCount = lifecycle::deprecated()
)
```

#### **Arguments**

```
x Vector of concept IDscdm cdm_reference via CDMConnector::cdm_from_con()
```

summariseCohortCodeUse 27

cohortTable A cohort table from the cdm reference.

cohortId A vector of cohort IDs to include

timing When to assess the code use relative cohort dates. This can be "any" (code use

any time by individuals in the cohort) or "entry" (code use on individuals' cohort

start date).

countBy Either "record" for record-level counts or "person" for person-level counts

byConcept TRUE or FALSE. If TRUE code use will be summarised by

by Year TRUE or FALSE. If TRUE code use will be summarised by year.

bySex TRUE or FALSE. If TRUE code use will be summarised by sex.

ageGroup If not NULL, a list of ageGroup vectors of length two.

 $\label{local-count} win Cell Count $$ \if else { html} {\href{https://lifecycle.r-lib.org/articles/stages.html $$ \else { html } {\href{https://lifecycle.r-lib.org/articles/stages.html $$ \else { html }$ \else { html }$$ 

#### Value

A tibble with results overall and, if specified, by strata

#### **Examples**

```
## Not run:
con <- DBI::dbConnect(duckdb::duckdb(),</pre>
                       dbdir = CDMConnector::eunomia_dir())
cdm <- CDMConnector::cdm_from_con(con,</pre>
                                   cdm_schem = "main",
                                   write_schema = "main")
cdm <- CDMConnector::generateConceptCohortSet(cdm = cdm,</pre>
conceptSet = list(a = 260139,
                  b = 1127433),
                  name = "cohorts",
                  end = "observation_period_end_date",
                  overwrite = TRUE)
results_cohort_mult <-
summariseCohortCodeUse(list(cs = c(260139,19133873)),
                       cdm = cdm,
                       cohortTable = "cohorts",
                       timing = "entry")
results_cohort_mult
CDMConnector::cdmDisconnect(cdm)
## End(Not run)
```

summariseOrphanCodes

Find orphan codes related to a codelist using achilles counts and, if available, PHOEBE concept recommendations

#### **Description**

Find orphan codes related to a codelist using achilles counts and, if available, PHOEBE concept recommendations

#### Usage

```
summariseOrphanCodes(
    x,
    cdm,
    domain = c("condition", "device", "drug", "measurement", "observation", "procedure",
        "visit")
)
```

#### **Arguments**

x A codelist for which to find related codes used in the database

cdm\_reference via CDMConnector

domain The domains to restrict results too. Only concepts from these domains will be

returned.

#### Value

A summarised result containg the frequency of codes related to (but not in) the codelist

#### **Examples**

```
cdm <- mockVocabRef("database")
codes <- getCandidateCodes(cdm = cdm,
keywords = "Musculoskeletal disorder",
domains = "Condition",
includeDescendants = FALSE)

orphan_codes <- summariseOrphanCodes(x = list("msk" = codes$concept_id),
cdm = cdm)

orphan_codes
CDMConnector::cdmDisconnect(cdm)</pre>
```

tableAchillesCodeUse 29

tableAchillesCodeUse Format the result of summariseAchillesCodeUse into a table.

#### **Description**

Format the result of summariseAchillesCodeUse into a table.

#### Usage

```
tableAchillesCodeUse(
  result,
  type = "gt",
  header = c("cdm_name", "estimate_name"),
  groupColumns = character(),
  hide = character(),
  .options = list(),
  conceptId = lifecycle::deprecated(),
  standard = lifecycle::deprecated(),
  vocabulary = lifecycle::deprecated(),
  excludeColumns = lifecycle::deprecated())
```

#### **Arguments**

result	A <summarised_res< th=""><th>sult&gt; with results o</th><th>f the type</th><th>"achilles code</th><th>use".</th></summarised_res<>	sult> with results o	f the type	"achilles code	use".

type Type of desired formatted table. To see supported formats use visOmopRe-

sults::tableType()

header A vector specifying the elements to include in the header. The order of el-

ements matters, with the first being the topmost header. The header vector can contain one of the following variables: "cdm\_name", "codelist\_name", "domain\_id", "standard\_concept\_name", "standard\_concept\_id", "estimate\_name", "standard\_concept", "vocabulary\_id". Alternatively, it can include other names

to use as overall header labels.

groupColumns Variables to use as group labels. Allowed columns are: "cdm\_name", "codelist\_name",

"domain\_id", "standard\_concept\_name", "standard\_concept\_id", "estimate\_name",

"standard\_concept", "vocabulary\_id". These cannot be used in header.

hide Table columns to exclude, options are: "cdm\_name", "codelist\_name", "do-

main\_id", "standard\_concept\_name", "standard\_concept\_id", "estimate\_name", "standard\_concept", "vocabulary\_id". These cannot be used in header or group-

Column.

. options Named list with additional formatting options. visOmopResults::tableOptions()

shows allowed arguments and their default values.

conceptId Deprecated. standard Deprecated. vocabulary Deprecated. excludeColumns Deprecated.

30 tableCodeUse

#### Value

A table with a formatted version of the summariseCohortCodeUse result.

#### **Examples**

```
cdm <- mockVocabRef("database")
oa <- getCandidateCodes(cdm = cdm, keywords = "osteoarthritis")
result_achilles <- summariseAchillesCodeUse(list(oa = oa$concept_id), cdm = cdm)
tableAchillesCodeUse(result_achilles)
CDMConnector::cdmDisconnect(cdm)</pre>
```

tableCodeUse

Format the result of summariseCodeUse into a table.

### **Description**

Format the result of summariseCodeUse into a table.

#### Usage

```
tableCodeUse(
  result,
  type = "gt",
  header = c("cdm_name", "estimate_name"),
  groupColumns = character(),
  hide = character(),
  .options = list(),
  splitStrata = lifecycle::deprecated(),
  conceptId = lifecycle::deprecated(),
  sourceConcept = lifecycle::deprecated(),
  excludeColumns = lifecycle::deprecated())
```

#### **Arguments**

result A summarised result with results of the type "code\_use".

type Type of desired formatted table. To see supported formats use visOmopRe-

sults::tableType()

header A vector specifying the elements to include in the header. The order of elements matters, with the first being the topmost header. The header vector can

contain one of the following variables: "cdm\_name", "codelist\_name", "standard\_concept\_name", "standard\_concept\_id", "estimate\_name", "source\_concept\_name", "source\_concept\_id", "domain\_id". If results are stratified, "year", "sex", "age\_group" can also be used. Alternatively, it can include other names to use as overall

header labels.

tableCohortCodeUse 31

groupColumns Variables to use as group labels. Allowed columns are: "cdm\_name", "codelist\_name",

"standard\_concept\_name", "standard\_concept\_id", "estimate\_name", "source\_concept\_name",

"source\_concept\_id", "domain\_id". If results are stratified, "year", "sex", "age\_group"

can also be used. These cannot be used in header.

hide Table columns to exclude, options are: "cdm\_name", "codelist\_name", "year",

"sex", "age\_group", "standard\_concept\_name", "standard\_concept\_id", "estimate\_name", "source\_concept\_name", "source\_concept\_id", "domain\_id". If results are stratified, "year", "sex", "age\_group" can also be used. These cannot

be used in header or groupColumn.

. options Named list with additional formatting options. visOmopResults::tableOptions()

shows allowed arguments and their default values.

splitStrata Deprecated.
conceptId Deprecated.
sourceConcept Deprecated.
excludeColumns Deprecated.

#### Value

A table with a formatted version of the summariseCodeUse result.

#### **Examples**

tableCohortCodeUse

Format the result of summariseCohortCodeUse into a table.

#### **Description**

Format the result of summariseCohortCodeUse into a table.

32 tableCohortCodeUse

#### Usage

```
tableCohortCodeUse(
  result,
  type = "gt",
  header = c("cdm_name", "estimate_name"),
  groupColumns = NULL,
  timing = FALSE,
  hide = character(),
  .options = list(),
  excludeColumns = lifecycle::deprecated(),
  splitStrata = lifecycle::deprecated(),
  conceptId = lifecycle::deprecated(),
  sourceConcept = lifecycle::deprecated())
```

#### **Arguments**

result A summarised result with results of the type "cohort\_code\_use".

type Type of desired formatted table. To see supported formats use visOmopRe-

sults::tableType()

header A vector specifying the elements to include in the header. The order of ele-

ments matters, with the first being the topmost header. The header vector can contain one of the following variables: "cdm\_name", "codelist\_name", "stan-

dard\_concept\_name", "standard\_concept\_id", "estimate\_name", "source\_concept\_name", "source\_concept\_id", "domain\_id". If results are stratified, "year", "sex", "age\_group" can also be used. Alternatively, it can include other names to use as overall

header labels.

groupColumns Variables to use as group labels. Allowed columns are: "cdm name", "codelist name",

"standard\_concept\_name", "standard\_concept\_id", "estimate\_name", "source\_concept\_name",

"source\_concept\_id", "domain\_id". If results are stratified, "year", "sex", "age\_group"

can also be used. These cannot be used in header.

timing If TRUE the timing setting will be displayed.

hide Table columns to exclude, options are: "cdm\_name", "codelist\_name", "year",

"sex", "age\_group", "standard\_concept\_name", "standard\_concept\_id", "estimate\_name", "source\_concept\_name", "source\_concept\_id", "domain\_id". If results are stratified, "year", "sex", "age\_group" can also be used. These cannot

be used in header or groupColumn.

options Named list with additional formatting options. visOmopResults::tableOptions()

shows allowed arguments and their default values.

excludeColumns Deprecated.
splitStrata Deprecated.
conceptId Deprecated.
sourceConcept Deprecated.

#### Value

A table with a formatted version of the summariseCohortCodeUse result.

tableOrphanCodes 33

#### **Examples**

```
con <- DBI::dbConnect(duckdb::duckdb(),</pre>
                       dbdir = CDMConnector::eunomia_dir())
cdm <- CDMConnector::cdm_from_con(con,</pre>
                                   cdm_schem = "main",
                                   write_schema = "main")
cdm <- CDMConnector::generateConceptCohortSet(cdm = cdm,</pre>
conceptSet = list(a = 260139,
                  b = 1127433),
                  name = "cohorts",
                   end = "observation_period_end_date",
                   overwrite = TRUE)
results_cohort_mult <-
summariseCohortCodeUse(list(cs = c(260139,19133873)),
                       cdm = cdm,
                       cohortTable = "cohorts",
                       timing = "entry")
tableCohortCodeUse(results_cohort_mult)
CDMConnector::cdmDisconnect(cdm)
## End(Not run)
```

tableOrphanCodes

Format the result of summariseOrphanCodes into a table.

#### **Description**

Format the result of summariseOrphanCodes into a table.

#### Usage

```
tableOrphanCodes(
  result,
  type = "gt",
  header = c("cdm_name", "estimate_name"),
  groupColumns = character(),
  hide = character(),
  .options = list(),
  conceptId = lifecycle::deprecated(),
  standard = lifecycle::deprecated(),
  vocabulary = lifecycle::deprecated(),
  excludeColumns = lifecycle::deprecated())
```

34 tableOrphanCodes

#### **Arguments**

result A summarised result with results of the type "orphan\_codes".

type Type of desired formatted table. To see supported formats use visOmopRe-

sults::tableType()

header A vector specifying the elements to include in the header. The order of el-

ements matters, with the first being the topmost header. The header vector can contain one of the following variables: "cdm\_name", "codelist\_name", "domain\_id", "standard\_concept\_name", "standard\_concept\_id", "estimate\_name", "standard\_concept", "vocabulary\_id". Alternatively, it can include other names

to use as overall header labels.

groupColumns Variables to use as group labels. Allowed columns are: "cdm\_name", "codelist\_name",

"domain\_id", "standard\_concept\_name", "standard\_concept\_id", "estimate\_name",

"standard\_concept", "vocabulary\_id". These cannot be used in header.

hide Table columns to exclude, options are: "cdm\_name", "codelist\_name", "do-

main\_id", "standard\_concept\_name", "standard\_concept\_id", "estimate\_name", "standard\_concept", "vocabulary\_id". These cannot be used in header or group-

Column.

options Named list with additional formatting options. visOmopResults::tableOptions()

shows allowed arguments and their default values.

conceptId Deprecated.
standard Deprecated.
vocabulary Deprecated.
excludeColumns Deprecated.

#### Value

A table with a formatted version of the summariseOrphanCodes result.

#### **Examples**

```
cdm <- mockVocabRef("database")
codes <- getCandidateCodes(cdm = cdm,
keywords = "Musculoskeletal disorder",
domains = "Condition",
includeDescendants = FALSE)

orphan_codes <- summariseOrphanCodes(x = list("msk" = codes$concept_id),
cdm = cdm)

tableOrphanCodes(orphan_codes)

CDMConnector::cdmDisconnect(cdm)</pre>
```

# **Index**

* datasets doseFormToRoute, 7
availableATC, 3
availableICD10,3
availableIngredients,4
codesFromCohort, 4
codesFromConceptSet, 5
codesInUse, 5
compareCodelists,6
doseFormToRoute, 7
getATCCodes, 7
${\tt getCandidateCodes}, 8$
getConceptClassId, 10
getDescendants, 10
getDomains, 11
getDoseForm, 12
getDoseUnit, 13
getDrugIngredientCodes, 13
getICD10StandardCodes, 14 getMappings, 15
getRelationshipId, 16
getRouteCategories, 17
getVocabularies, 18
getVocabVersion, 18
mockVocabRef, 19
restrictToCodesInUse, 19
sourceCodesInUse, 20
stratifyByConcept, 21
stratifyByDoseUnit, 21
stratifyByRouteCategory, 22
subsetOnDoseUnit, 22
subsetOnRouteCategory, 23
subsetToCodesInUse, 23
summariseAchillesCodeUse, 24

```
summariseCodeUse, 25
summariseCohortCodeUse, 26
summariseOrphanCodes, 28
table A chilles Code Use, {\color{red}29}
tableCodeUse, 30
tableCohortCodeUse, 31
tableOrphanCodes, 33
```