

Package ‘DrugUtilisation’

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Title Summarise Patient-Level Drug Utilisation in Data Mapped to the OMOP Common Data Model

Version 1.0.4

Description Summarise patient-level drug utilisation cohorts using data mapped to the Observational Medical Outcomes Partnership (OMOP) common data model. New users and prevalent users cohorts can be generated and their characteristics, indication and drug use summarised.

License Apache License (>= 2)

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Imports CDMConnector (>= 1.4.0), cli, clock, CodelistGenerator (>= 3.1.0), dplyr, glue, omopgenerics (>= 1.0.0), PatientProfiles (>= 1.0.0), purrr, rlang, stringr, tidyverse

Depends R (>= 4.1)

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URL <https://darwin-eu.github.io/DrugUtilisation/>

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addCumulativeDose	<i>To add a new column with the cumulative dose. To add multiple columns use addDrugUtilisation() for efficiency.</i>
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Description

To add a new column with the cumulative dose. To add multiple columns use `addDrugUtilisation()` for efficiency.

Usage

```
addCumulativeDose(  
  cohort,  
  ingredientConceptId,  
  conceptSet = NULL,  
  indexDate = "cohort_start_date",  
  censorDate = "cohort_end_date",  
  restrictIncident = TRUE,  
  nameStyle = "cumulative_dose_{concept_name}_{ingredient}",  
  name = NULL  
)
```

Arguments

<code>cohort</code>	A cohort_table object.
<code>ingredientConceptId</code>	Ingredient OMOP concept that we are interested for the study.
<code>conceptSet</code>	List of concepts to be included.
<code>indexDate</code>	Name of a column that indicates the date to start the analysis.
<code>censorDate</code>	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
<code>restrictIncident</code>	Whether to include only incident prescriptions in the analysis. If FALSE all prescriptions that overlap with the study period will be included.
<code>nameStyle</code>	Character string to specify the nameStyle of the new columns.

<code>name</code>	Name of the new computed cohort table, if NULL a temporary table will be created.
-------------------	---

Value

The same cohort with the added column.

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()
codelist <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "dus_cohort",
                                         conceptSet = codelist)

cdm$dus_cohort |>
  addCumulativeDose(ingredientConceptId = 1125315) |>
  glimpse()
```

`addCumulativeQuantity` *To add a new column with the cumulative quantity. To add multiple columns use addDrugUtilisation() for efficiency.*

Description

To add a new column with the cumulative quantity. To add multiple columns use `addDrugUtilisation()` for efficiency.

Usage

```
addCumulativeQuantity(
  cohort,
  conceptSet,
  indexDate = "cohort_start_date",
  censorDate = "cohort_end_date",
  restrictIncident = TRUE,
  nameStyle = "cumulative_quantity_{concept_name}",
  name = NULL
)
```

Arguments

cohort	A cohort_table object.
conceptSet	List of concepts to be included.
indexDate	Name of a column that indicates the date to start the analysis.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
restrictIncident	Whether to include only incident prescriptions in the analysis. If FALSE all prescriptions that overlap with the study period will be included.
nameStyle	Character string to specify the nameStyle of the new columns.
name	Name of the new computed cohort table, if NULL a temporary table will be created.

Value

The same cohort with the added column.

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()
codelist <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "dus_cohort",
                                         conceptSet = codelist)

cdm$dus_cohort |>
  addCumulativeQuantity(conceptSet = codelist) |>
  glimpse()
```

`addDaysExposed`

To add a new column with the days exposed. To add multiple columns use addDrugUtilisation() for efficiency.

Description

To add a new column with the days exposed. To add multiple columns use `addDrugUtilisation()` for efficiency.

Usage

```
addDaysExposed(
  cohort,
  conceptSet,
  gapEra,
  indexDate = "cohort_start_date",
  censorDate = "cohort_end_date",
  restrictIncident = TRUE,
  nameStyle = "days_exposed_{concept_name}",
  name = NULL
)
```

Arguments

cohort	A cohort_table object.
conceptSet	List of concepts to be included.
gapEra	Number of days between two continuous exposures to be considered in the same era.
indexDate	Name of a column that indicates the date to start the analysis.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
restrictIncident	Whether to include only incident prescriptions in the analysis. If FALSE all prescriptions that overlap with the study period will be included.
nameStyle	Character string to specify the nameStyle of the new columns.
name	Name of the new computed cohort table, if NULL a temporary table will be created.

Value

The same cohort with the added column.

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()
codelist <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "dus_cohort",
                                         conceptSet = codelist)

cdm$dus_cohort |>
  addDaysExposed(conceptSet = codelist, gapEra = 1) |>
  glimpse()
```

addDaysPrescribed *To add a new column with the days prescribed. To add multiple columns use addDrugUtilisation() for efficiency.*

Description

To add a new column with the days prescribed. To add multiple columns use addDrugUtilisation() for efficiency.

Usage

```
addDaysPrescribed(  
  cohort,  
  conceptSet,  
  indexDate = "cohort_start_date",  
  censorDate = "cohort_end_date",  
  restrictIncident = TRUE,  
  nameStyle = "days_prescribed_{concept_name}",  
  name = NULL  
)
```

Arguments

cohort	A cohort_table object.
conceptSet	List of concepts to be included.
indexDate	Name of a column that indicates the date to start the analysis.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
restrictIncident	Whether to include only incident prescriptions in the analysis. If FALSE all prescriptions that overlap with the study period will be included.
nameStyle	Character string to specify the nameStyle of the new columns.
name	Name of the new computed cohort table, if NULL a temporary table will be created.

Value

The same cohort with the added columns.

Examples

```
library(DrugUtilisation)  
library(CodelistGenerator)  
library(dplyr, warn.conflicts = FALSE)  
  
cdm <- mockDrugUtilisation()
```

```

codelist <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "dus_cohort",
                                         conceptSet = codelist)

cdm$dus_cohort |>
  addDaysPrescribed(conceptSet = codelist) |>
  glimpse()

```

addDrugRestart

Add drug restart information as a column per follow-up period of interest.

Description

Add drug restart information as a column per follow-up period of interest.

Usage

```

addDrugRestart(
  cohort,
  switchCohortTable,
  switchCohortId = NULL,
  followUpDays = Inf,
  censorDate = NULL,
  incident = TRUE,
  nameStyle = "drug_restart_{follow_up_days}"
)

```

Arguments

- cohort** A cohort_table object.
- switchCohortTable** A cohort table in the cdm that contains possible alternative treatments.
- switchCohortId** The cohort ids to be used from switchCohortTable. If NULL all cohort definition ids are used.
- followUpDays** A vector of number of days to follow up. It can be multiple values.
- censorDate** Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
- incident** Whether the switch treatment has to be incident (start after discontinuation) or not (it can start before the discontinuation and last till after).
- nameStyle** Character string to specify the nameStyle of the new columns.

Value

The cohort table given with additional columns with information on the restart, switch and not exposed per follow-up period of interest.

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation()

conceptlist <- list(acetaminophen = 1125360, metformin = c(1503297, 1503327))
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "switch_cohort",
                                         conceptSet = conceptlist)

cdm$cohort1 |>
  addDrugRestart(switchCohortTable = "switch_cohort")
```

addDrugUtilisation *Add new columns with drug use related information*

Description

Add new columns with drug use related information

Usage

```
addDrugUtilisation(
  cohort,
  gapEra,
  conceptSet = NULL,
  ingredientConceptId = NULL,
  indexDate = "cohort_start_date",
  censorDate = "cohort_end_date",
  restrictIncident = TRUE,
  numberExposures = TRUE,
  numberEras = TRUE,
  daysExposed = TRUE,
  daysPrescribed = TRUE,
  timeToExposure = TRUE,
  initialExposureDuration = TRUE,
  initialQuantity = TRUE,
  cumulativeQuantity = TRUE,
  initialDailyDose = TRUE,
  cumulativeDose = TRUE,
  nameStyle = "{value}_{concept_name}_{ingredient}",
  name = NULL
)
```

Arguments

cohort	A cohort_table object.
gapEra	Number of days between two continuous exposures to be considered in the same era.
conceptSet	List of concepts to be included.
ingredientConceptId	Ingredient OMOP concept that we are interested for the study.
indexDate	Name of a column that indicates the date to start the analysis.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
restrictIncident	Whether to include only incident prescriptions in the analysis. If FALSE all prescriptions that overlap with the study period will be included.
numberExposures	Whether to include 'number_exposures' (number of drug exposure records between indexDate and censorDate).
numberEras	Whether to include 'number_eras' (number of continuous exposure episodes between indexDate and censorDate).
daysExposed	Whether to include 'days_exposed' (number of days that the individual is in a continuous exposure episode, including allowed treatment gaps, between indexDate and censorDate; sum of the length of the different drug eras).
daysPrescribed	Whether to include 'days_prescribed' (sum of the number of days for each prescription that contribute in the analysis).
timeToExposure	Whether to include 'time_to_exposure' (number of days between indexDate and the first episode).
initialExposureDuration	Whether to include 'initial_exposure_duration' (number of prescribed days of the first drug exposure record).
initialQuantity	Whether to include 'initial_quantity' (quantity of the first drug exposure record).
cumulativeQuantity	Whether to include 'cumulative_quantity' (sum of the quantity of the different exposures considered in the analysis).
initialDailyDose	Whether to include 'initial_daily_dose_{unit}' (daily dose of the first considered prescription).
cumulativeDose	Whether to include 'cumulative_dose_{unit}' (sum of the cumulative dose of the analysed drug exposure records).
nameStyle	Character string to specify the nameStyle of the new columns.
name	Name of the new computed cohort table, if NULL a temporary table will be created.

Value

The same cohort with the added columns.

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()
codelist <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")

cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "dus_cohort",
                                         conceptSet = codelist)
cdm$dus_cohort |>
  addDrugUtilisation(ingredientConceptId = 1125315, gapEra = 30) |>
  glimpse()
```

addIndication

Add a variable indicating individuals indications

Description

Add a variable to a drug cohort indicating their presence in an indication cohort in a specified time window. If an individual is not in one of the indication cohorts, they will be considered to have an unknown indication if they are present in one of the specified OMOP CDM clinical tables. If they are neither in an indication cohort or a clinical table they will be considered as having no observed indication.

Usage

```
addIndication(
  cohort,
  indicationCohortName,
  indicationCohortId = NULL,
  indicationWindow = list(c(0, 0)),
  unknownIndicationTable = NULL,
  indexDate = "cohort_start_date",
  censorDate = NULL,
  mutuallyExclusive = TRUE,
  nameStyle = NULL,
  name = NULL
)
```

Arguments

cohort	A cohort_table object.
indicationCohortName	Name of indication cohort table

indicationCohortId	target cohort Id to add indication
indicationWindow	time window of interests
unknownIndicationTable	Tables to search unknown indications
indexDate	Name of a column that indicates the date to start the analysis.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
mutuallyExclusive	Whether to consider mutually exclusive categories (one column per window) or not (one column per window and indication).
nameStyle	Name style for the indications. By default: 'indication_{window_name}' (mutuallyExclusive = TRUE), 'indication_{window_name}_{cohort_name}' (mutuallyExclusive = FALSE).
name	Name of the new computed cohort table, if NULL a temporary table will be created.

Value

The original table with a variable added that summarises the individual's indications.

Examples

```
library(DrugUtilisation)
library(CDMConnector)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()

indications <- list(headache = 378253, asthma = 317009)
cdm <- generateConceptCohortSet(cdm = cdm,
                                  conceptSet = indications,
                                  name = "indication_cohorts")

cdm <- generateIngredientCohortSet(cdm = cdm,
                                      name = "drug_cohort",
                                      ingredient = "acetaminophen")

cdm$drug_cohort |>
  addIndication(
    indicationCohortName = "indication_cohorts",
    indicationWindow = list(c(0, 0)),
    unknownIndicationTable = "condition_occurrence"
  ) |>
  glimpse()
```

addInitialDailyDose *To add a new column with the initial daily dose. To add multiple columns use addDrugUtilisation() for efficiency.*

Description

To add a new column with the initial daily dose. To add multiple columns use addDrugUtilisation() for efficiency.

Usage

```
addInitialDailyDose(  
  cohort,  
  ingredientConceptId,  
  conceptSet = NULL,  
  indexDate = "cohort_start_date",  
  censorDate = "cohort_end_date",  
  restrictIncident = TRUE,  
  nameStyle = "initial_daily_dose_{concept_name}_{ingredient}",  
  name = NULL  
)
```

Arguments

cohort	A cohort_table object.
ingredientConceptId	Ingredient OMOP concept that we are interested for the study.
conceptSet	List of concepts to be included.
indexDate	Name of a column that indicates the date to start the analysis.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
restrictIncident	Whether to include only incident prescriptions in the analysis. If FALSE all prescriptions that overlap with the study period will be included.
nameStyle	Character string to specify the nameStyle of the new columns.
name	Name of the new computed cohort table, if NULL a temporary table will be created.

Value

The same cohort with the added column.

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()
codelist <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "dus_cohort",
                                         conceptSet = codelist)

cdm$dus_cohort |>
  addInitialDailyDose(ingredientConceptId = 1125315) |>
  glimpse()
```

`addInitialExposureDuration`

To add a new column with the duration of the first exposure. To add multiple columns use `addDrugUtilisation()` for efficiency.

Description

To add a new column with the duration of the first exposure. To add multiple columns use `addDrugUtilisation()` for efficiency.

Usage

```
addInitialExposureDuration(
  cohort,
  conceptSet,
  indexDate = "cohort_start_date",
  censorDate = "cohort_end_date",
  restrictIncident = TRUE,
  nameStyle = "initial_exposure_duration_{concept_name}",
  name = NULL
)
```

Arguments

<code>cohort</code>	A cohort_table object.
<code>conceptSet</code>	List of concepts to be included.
<code>indexDate</code>	Name of a column that indicates the date to start the analysis.
<code>censorDate</code>	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.

<code>restrictIncident</code>	Whether to include only incident prescriptions in the analysis. If FALSE all prescriptions that overlap with the study period will be included.
<code>nameStyle</code>	Character string to specify the nameStyle of the new columns.
<code>name</code>	Name of the new computed cohort table, if NULL a temporary table will be created.

Value

The same cohort with the added column.

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()
codelist <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "dus_cohort",
                                         conceptSet = codelist)

cdm$dus_cohort |>
  addInitialExposureDuration(conceptSet = codelist) |>
  glimpse()
```

`addInitialQuantity` *To add a new column with the initial quantity. To add multiple columns use addDrugUtilisation() for efficiency.*

Description

To add a new column with the initial quantity. To add multiple columns use `addDrugUtilisation()` for efficiency.

Usage

```
addInitialQuantity(
  cohort,
  conceptSet,
  indexDate = "cohort_start_date",
  censorDate = "cohort_end_date",
  restrictIncident = TRUE,
  nameStyle = "initial_quantity_{concept_name}",
  name = NULL
)
```

Arguments

<code>cohort</code>	A cohort_table object.
<code>conceptSet</code>	List of concepts to be included.
<code>indexDate</code>	Name of a column that indicates the date to start the analysis.
<code>censorDate</code>	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
<code>restrictIncident</code>	Whether to include only incident prescriptions in the analysis. If FALSE all prescriptions that overlap with the study period will be included.
<code>nameStyle</code>	Character string to specify the nameStyle of the new columns.
<code>name</code>	Name of the new computed cohort table, if NULL a temporary table will be created.

Value

The same cohort with the added column.

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()
codelist <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "dus_cohort",
                                         conceptSet = codelist)

cdm$dus_cohort |>
  addInitialQuantity(conceptSet = codelist) |>
  glimpse()
```

<code>addNumberEras</code>	<i>To add a new column with the number of eras. To add multiple columns use addDrugUtilisation() for efficiency.</i>
----------------------------	--

Description

To add a new column with the number of eras. To add multiple columns use `addDrugUtilisation()` for efficiency.

Usage

```
addNumberEras(
  cohort,
  conceptSet,
  gapEra,
  indexDate = "cohort_start_date",
  censorDate = "cohort_end_date",
  restrictIncident = TRUE,
  nameStyle = "number_eras_{concept_name}",
  name = NULL
)
```

Arguments

cohort	A cohort_table object.
conceptSet	List of concepts to be included.
gapEra	Number of days between two continuous exposures to be considered in the same era.
indexDate	Name of a column that indicates the date to start the analysis.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
restrictIncident	Whether to include only incident prescriptions in the analysis. If FALSE all prescriptions that overlap with the study period will be included.
nameStyle	Character string to specify the nameStyle of the new columns.
name	Name of the new computed cohort table, if NULL a temporary table will be created.

Value

The same cohort with the added column.

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()
codelist <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "dus_cohort",
                                         conceptSet = codelist)

cdm$dus_cohort |>
  addNumberEras(conceptSet = codelist, gapEra = 1) |>
  glimpse()
```

addNumberExposures	<i>To add a new column with the number of exposures. To add multiple columns use addDrugUtilisation() for efficiency.</i>
--------------------	---

Description

To add a new column with the number of exposures. To add multiple columns use addDrugUtilisation() for efficiency.

Usage

```
addNumberExposures(
  cohort,
  conceptSet,
  indexDate = "cohort_start_date",
  censorDate = "cohort_end_date",
  restrictIncident = TRUE,
  nameStyle = "number_exposures_{concept_name}",
  name = NULL
)
```

Arguments

cohort	A cohort_table object.
conceptSet	List of concepts to be included.
indexDate	Name of a column that indicates the date to start the analysis.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
restrictIncident	Whether to include only incident prescriptions in the analysis. If FALSE all prescriptions that overlap with the study period will be included.
nameStyle	Character string to specify the nameStyle of the new columns.
name	Name of the new computed cohort table, if NULL a temporary table will be created.

Value

The same cohort with the added columns.

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()
```

```

cdelist <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "dus_cohort",
                                         conceptSet = codelist)

cdm$dus_cohort |>
  addNumberExposures(conceptSet = codelist) |>
  glimpse()

```

addTimeToExposure

To add a new column with the time to exposure. To add multiple columns use addDrugUtilisation() for efficiency.

Description

To add a new column with the time to exposure. To add multiple columns use addDrugUtilisation() for efficiency.

Usage

```

addTimeToExposure(
  cohort,
  conceptSet,
  indexDate = "cohort_start_date",
  censorDate = "cohort_end_date",
  restrictIncident = TRUE,
  nameStyle = "time_to_exposure_{concept_name}",
  name = NULL
)

```

Arguments

<code>cohort</code>	A cohort_table object.
<code>conceptSet</code>	List of concepts to be included.
<code>indexDate</code>	Name of a column that indicates the date to start the analysis.
<code>censorDate</code>	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
<code>restrictIncident</code>	Whether to include only incident prescriptions in the analysis. If FALSE all prescriptions that overlap with the study period will be included.
<code>nameStyle</code>	Character string to specify the nameStyle of the new columns.
<code>name</code>	Name of the new computed cohort table, if NULL a temporary table will be created.

Value

The same cohort with the added column.

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()
codelist <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "dus_cohort",
                                         conceptSet = codelist)

cdm$dus_cohort |>
  addTimeToExposure(conceptSet = codelist) |>
  glimpse()
```

addTreatment

Add a variable indicating individuals medications

Description

Add a variable to a drug cohort indicating their presence of a medication cohort in a specified time window.

Usage

```
addTreatment(
  cohort,
  treatmentCohortName,
  treatmentCohortId = NULL,
  window = list(c(0, 0)),
  indexDate = "cohort_start_date",
  censorDate = NULL,
  mutuallyExclusive = TRUE,
  nameStyle = NULL,
  name = NULL
)
```

Arguments

cohort	A cohort_table object.
treatmentCohortName	Name of treatment cohort table

treatmentCohortId	target cohort Id to add treatment
window	time window of interests.
indexDate	Name of a column that indicates the date to start the analysis.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
mutuallyExclusive	Whether to consider mutually exclusive categories (one column per window) or not (one column per window and treatment).
nameStyle	Name style for the treatment columns. By default: 'treatment_{window_name}' (mutuallyExclusive = TRUE), 'treatment_{window_name}_{cohort_name}' (mutuallyExclusive = FALSE).
name	Name of the new computed cohort table, if NULL a temporary table will be created.

Value

The original table with a variable added that summarises the individual's indications.

Examples

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

cdm <- mockDrugUtilisation(numberIndividuals = 50)

cdm <- generateIngredientCohortSet(cdm = cdm,
                                      name = "drug_cohort",
                                      ingredient = "acetaminophen")

cdm <- generateIngredientCohortSet(cdm = cdm,
                                      name = "treatments",
                                      ingredient = c("metformin", "simvastatin"))

cdm$drug_cohort |>
  addTreatment("treatments", window = list(c(0, 0), c(1, 30), c(31, 60))) |>
  glimpse()
```

Description

Run benchmark of drug utilisation cohort generation

Usage

```
benchmarkDrugUtilisation(
  cdm,
  ingredient = "acetaminophen",
  alternativeIngredient = c("ibuprofen", "aspirin", "diclofenac"),
  indicationCohort = NULL
)
```

Arguments

cdm	A cdm_reference object.
ingredient	Name of ingredient to benchmark.
alternativeIngredient	Name of ingredients to use as alternative treatments.
indicationCohort	Name of a cohort in the cdm_reference object to use as indicationm.

Value

A summarise_result object.

Examples

```
library(DrugUtilisation)
library(CDMConnector)
library(duckdb)

requireEunomia()
con <- dbConnect(drv = duckdb(dbdir = eunomiaDir()))
cdm <- cdmFromCon(con = con, cdmSchema = "main", writeSchema = "main")

timings <- benchmarkDrugUtilisation(cdm)

timings
```

cohortGapEra

Get the gapEra used to create a cohort

Description

Get the gapEra used to create a cohort

Usage

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

Arguments

- cohort A cohort_table object.
- cohortId Integer vector referring to cohortIds from cohort. If NULL all cohort definition ids in settings will be used.

Value

gapEra values for the specific cohortIds

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)

cdm <- mockDrugUtilisation()

druglist <- getDrugIngredientCodes(cdm = cdm,
                                     name = c("acetaminophen", "metformin"))

cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                           name = "drug_cohorts",
                                           conceptSet = druglist,
                                           gapEra = 100)

cohortGapEra(cdm$drug_cohorts)
```

erafyCohort

Erafy a cohort_table collapsing records separated gapEra days or less.

Description

Erafy a cohort_table collapsing records separated gapEra days or less.

Usage

```
erafyCohort(
  cohort,
  gapEra,
  cohortId = NULL,
  nameStyle = "{cohort_name}_{gap_era}",
  name = omopgenerics::tableName(cohort)
)
```

Arguments

cohort	A cohort_table object.
gapEra	Number of days between two continuous exposures to be considered in the same era.
cohortId	A cohort definition id to restrict by. If NULL, all cohorts will be included.
nameStyle	String to create the new names of cohorts. Must contain '{cohort_name}' if more than one cohort is present and '{gap_era}' if more than one gapEra is provided.
name	Name of the new cohort table, it must be a length 1 character vector.

Value

A cohort_table object.

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation()

cdm$cohort2 <- cdm$cohort1 |>
  erafyCohort(gapEra = 30, name = "cohort2")

cdm$cohort2

settings(cdm$cohort2)

mockDisconnect(cdm)
```

generateAtcCohortSet *Generate a set of drug cohorts based on ATC classification*

Description

Adds a new cohort table to the cdm reference with individuals who have drug exposure records that belong to the specified Anatomical Therapeutic Chemical (ATC) classification. Cohort start and end dates will be based on drug record start and end dates, respectively. Records that overlap or have fewer days between them than the specified gap era will be concatenated into a single cohort entry.

Usage

```
generateAtcCohortSet(
  cdm,
  name,
  atcName = NULL,
```

```

gapEra = 1,
subsetCohort = NULL,
subsetCohortId = NULL,
numberExposures = FALSE,
daysPrescribed = FALSE,
...
)

```

Arguments

cdm	A cdm_reference object.
name	Name of the new cohort table, it must be a length 1 character vector.
atcName	Names of ATC classification of interest.
gapEra	Number of days between two continuous exposures to be considered in the same era. Records that have fewer days between them than this gap will be concatenated into the same cohort record.
subsetCohort	Cohort table to subset.
subsetCohortId	Cohort id to subset.
numberExposures	Whether to include 'number_exposures' (number of drug exposure records between indexDate and censorDate).
daysPrescribed	Whether to include 'days_prescribed' (number of days prescribed used to create each era).
...	Arguments to be passed to CodelistGenerator::getATCCodes().

Value

The function returns the cdm reference provided with the addition of the new cohort table.

Examples

```

library(DrugUtilisation)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()

cdm <- generateAtcCohortSet(cdm = cdm,
                             atcName = "alimentary tract and metabolism",
                             name = "drugs")

cdm$drugs |>
glimpse()

```

generateDrugUtilisationCohortSet

Generate a set of drug cohorts based on given concepts

Description

Adds a new cohort table to the cdm reference with individuals who have drug exposure records with the specified concepts. Cohort start and end dates will be based on drug record start and end dates, respectively. Records that overlap or have fewer days between them than the specified gap era will be concatenated into a single cohort entry.

Usage

```
generateDrugUtilisationCohortSet(
  cdm,
  name,
  conceptSet,
  gapEra = 1,
  subsetCohort = NULL,
  subsetCohortId = NULL,
  numberExposures = FALSE,
  daysPrescribed = FALSE
)
```

Arguments

cdm	A cdm_reference object.
name	Name of the new cohort table, it must be a length 1 character vector.
conceptSet	List of concepts to be included.
gapEra	Number of days between two continuous exposures to be considered in the same era.
subsetCohort	Cohort table to subset.
subsetCohortId	Cohort id to subset.
numberExposures	Whether to include 'number_exposures' (number of drug exposure records between indexDate and censorDate).
daysPrescribed	Whether to include 'days_prescribed' (number of days prescribed used to create each era).

Value

The function returns the cdm reference provided with the addition of the new cohort table.

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()

druglist <- getDrugIngredientCodes(cdm = cdm,
                                     name = c("acetaminophen", "metformin"),
                                     nameStyle = "{concept_name}")

cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                           name = "drug_cohorts",
                                           conceptSet = druglist,
                                           gapEra = 30,
                                           numberExposures = TRUE,
                                           daysPrescribed = TRUE)

cdm$drug_cohorts |>
  glimpse()
```

generateIngredientCohortSet

Generate a set of drug cohorts based on drug ingredients

Description

Adds a new cohort table to the cdm reference with individuals who have drug exposure records with the specified drug ingredient. Cohort start and end dates will be based on drug record start and end dates, respectively. Records that overlap or have fewer days between them than the specified gap era will be concatenated into a single cohort entry.

Usage

```
generateIngredientCohortSet(
  cdm,
  name,
  ingredient = NULL,
  gapEra = 1,
  subsetCohort = NULL,
  subsetCohortId = NULL,
  numberExposures = FALSE,
  daysPrescribed = FALSE,
  ...
)
```

Arguments

<code>cdm</code>	A <code>cdm_reference</code> object.
<code>name</code>	Name of the new cohort table, it must be a length 1 character vector.
<code>ingredient</code>	Accepts both vectors and named lists of ingredient names. For a vector input, e.g., <code>c("acetaminophen", "codeine")</code> , it generates a cohort table with descendant concept codes for each ingredient, assigning unique <code>cohort_definition_id</code> . For a named list input, e.g., <code>list("test_1" = c("simvastatin", "acetaminophen"), "test_2" = "metformin")</code> , it produces a cohort table based on the structure of the input, where each name leads to a combined set of descendant concept codes for the specified ingredients, creating distinct <code>cohort_definition_id</code> for each named group.
<code>gapEra</code>	Number of days between two continuous exposures to be considered in the same era.
<code>subsetCohort</code>	Cohort table to subset.
<code>subsetCohortId</code>	Cohort id to subset.
<code>numberExposures</code>	Whether to include 'number_exposures' (number of drug exposure records between <code>indexDate</code> and <code>censorDate</code>).
<code>daysPrescribed</code>	Whether to include 'days_prescribed' (number of days prescribed used to create each era).
<code>...</code>	Arguments to be passed to <code>CodelistGenerator::getDrugIngredientCodes()</code> .

Value

The function returns the `cdm` reference provided with the addition of the new cohort table.

Examples

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

cdm <- mockDrugUtilisation()

cdm <- generateIngredientCohortSet(cdm = cdm,
                                      ingredient = "acetaminophen",
                                      name = "acetaminophen")

cdm$acetaminophen |>
  glimpse()
```

mockDrugUtilisation *It creates a mock database for testing DrugUtilisation package*

Description

It creates a mock database for testing DrugUtilisation package

Usage

```
mockDrugUtilisation(  
  con = NULL,  
  writeSchema = NULL,  
  numberIndividuals = 10,  
  seed = NULL,  
  ...  
)
```

Arguments

con	A DBIConnection object to a database. If NULL a new duckdb connection will be used.
writeSchema	A schema with writing permissions to copy there the cdm tables.
numberIndividuals	Number of individuals in the mock cdm.
seed	Seed for the random numbers. If NULL no seed is used.
...	Tables to use as basis to create the mock. If some tables are provided they will be used to construct the cdm object.

Value

A cdm reference with the mock tables

Examples

```
library(DrugUtilisation)  
  
cdm <- mockDrugUtilisation()  
  
cdm
```

`patternsWithFormula` *Patterns valid to compute daily dose with the associated formula.*

Description

Patterns valid to compute daily dose with the associated formula.

Usage

```
patternsWithFormula
```

Format

A data frame with eight variables: pattern_id, amount, amount_unit, numerator, numerator_unit, denominator, denominator_unit, formula_name and formula.

`patternTable`

Function to create a tibble with the patterns from current drug strength table

Description

Function to create a tibble with the patterns from current drug strength table

Usage

```
patternTable(cdm)
```

Arguments

`cdm` A `cdm_reference` object.

Value

The function creates a tibble with the different patterns found in the table, plus a column of potentially valid and invalid combinations.

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation()

patternTable(cdm)
```

plotDrugRestart	<i>Generate a custom ggplot2 from a summarised_result object generated with summariseDrugRestart() function.</i>
-----------------	--

Description

Generate a custom ggplot2 from a summarised_result object generated with summariseDrugRestart() function.

Usage

```
plotDrugRestart(  
  result,  
  facet = cdm_name + cohort_name ~ follow_up_days,  
  colour = "variable_level"  
)
```

Arguments

result	A summarised_result object.
facet	Columns to facet by. See options with availablePlotColumns(result). Formula is also allowed to specify rows and columns.
colour	Columns to color by. See options with availablePlotColumns(result).

Value

A ggplot2 object.

Examples

```
## Not run:  
library(DrugUtilisation)  
  
cdm <- mockDrugUtilisation()  
  
conceptlist <- list("a" = 1125360, "b" = c(1503297, 1503327))  
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,  
                                         name = "switch_cohort",  
                                         conceptSet = conceptlist)  
  
result <- cdm$cohort1 |>  
         summariseDrugRestart(switchCohortTable = "switch_cohort")  
  
plotDrugRestart(result)  
  
## End(Not run)
```

`plotDrugUtilisation` *Plot the results of summariseDrugUtilisation*

Description

Plot the results of summariseDrugUtilisation

Usage

```
plotDrugUtilisation(
  result,
  variable = "number exposures",
  plotType = "barplot",
  facet = strataColumns(result),
  colour = "cohort_name"
)
```

Arguments

<code>result</code>	A summarised_result object.
<code>variable</code>	Variable to plot. See <code>unique(result\$variable_name)</code> for options.
<code>plotType</code>	Must be a choice between: 'scatterplot', 'barplot', 'densityplot', and 'boxplot'.
<code>facet</code>	Columns to facet by. See options with <code>availablePlotColumns(result)</code> . Formula is also allowed to specify rows and columns.
<code>colour</code>	Columns to color by. See options with <code>availablePlotColumns(result)</code> .

Value

A ggplot2 object.

Examples

```
library(DrugUtilisation)
library(PatientProfiles)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation(numberIndividuals = 100)
codes <- list(aceta = c(1125315, 1125360, 2905077, 43135274))
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "cohort",
                                         conceptSet = codes)

result <- cdm$cohort |>
  addSex() |>
  summariseDrugUtilisation(
    strata = "sex",
    ingredientConceptId = 1125315,
```

```
estimates = c("min", "q25", "median", "q75", "max", "density")
)

result |>
  filter(estimate_name == "median") |>
  plotDrugUtilisation(
    variable = "days prescribed",
    plotType = "barplot"
  )

result |>
  plotDrugUtilisation(
    variable = "days exposed",
    facet = cohort_name ~ cdm_name,
    colour = "sex",
    plotType = "boxplot"
  )

result |>
  plotDrugUtilisation(
    variable = "cumulative dose milligram",
    plotType = "densityplot",
    facet = "cohort_name",
    colour = "sex"
  )

mockDisconnect(cdm)
```

plotIndication

Generate a plot visualisation (ggplot2) from the output of summariseIndication

Description

Generate a plot visualisation (ggplot2) from the output of summariseIndication

Usage

```
plotIndication(
  result,
  facet = cdm_name + cohort_name ~ window_name,
  colour = "variable_level"
)
```

Arguments

result A summarised_result object.

<code>facet</code>	Columns to facet by. See options with <code>availablePlotColumns(result)</code> . Formula is also allowed to specify rows and columns.
<code>colour</code>	Columns to color by. See options with <code>availablePlotColumns(result)</code> .

Value

A ggplot2 object

Examples

```
library(DrugUtilisation)
library(CDMConnector)

cdm <- mockDrugUtilisation()

indications <- list(headache = 378253, asthma = 317009)
cdm <- generateConceptCohortSet(cdm = cdm,
                                  conceptSet = indications,
                                  name = "indication_cohorts")

cdm <- generateIngredientCohortSet(cdm = cdm,
                                      name = "drug_cohort",
                                      ingredient = "acetaminophen")

result <- cdm$drug_cohort |>
  summariseIndication(
    indicationCohortName = "indication_cohorts",
    unknownIndicationTable = "condition_occurrence",
    indicationWindow = list(c(-Inf, 0), c(-365, 0)))
  )

plotIndication(result)
```

plotProportionOfPatientsCovered

Plot proportion of patients covered

Description

Plot proportion of patients covered

Usage

```
plotProportionOfPatientsCovered(
  result,
  facet = "cohort_name",
  colour = strataColumns(result),
  ribbon = TRUE
)
```

Arguments

result	A summarised_result object.
facet	Columns to facet by. See options with availablePlotColumns(result). Formula is also allowed to specify rows and columns.
colour	Columns to color by. See options with availablePlotColumns(result).
ribbon	Whether to plot a ribbon with the confidence intervals.

Value

Plot of proportion Of patients covered over time

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation()

cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "my_cohort",
                                         conceptSet = list(drug_of_interest = c(1503297, 1503327)))

result <- cdm$my_cohort |>
  summariseProportionOfPatientsCovered(followUpDays = 365)

plotProportionOfPatientsCovered(result)
```

plotTreatment

Generate a custom ggplot2 from a summarised_result object generated with summariseTreatment function.

Description

Generate a custom ggplot2 from a summarised_result object generated with summariseTreatment function.

Usage

```
plotTreatment(
  result,
  facet = cdm_name + cohort_name ~ window_name,
  colour = "variable_level"
)
```

Arguments

<code>result</code>	A summarised_result object.
<code>facet</code>	Columns to facet by. See options with availablePlotColumns(result). Formula is also allowed to specify rows and columns.
<code>colour</code>	Columns to color by. See options with availablePlotColumns(result).

Value

A ggplot2 object.

Examples

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

cdm <- mockDrugUtilisation()
result <- cdm$cohort1 |>
  summariseTreatment(
    treatmentCohortName = "cohort2",
    window = list(c(0, 30), c(31, 365))
  )

plotTreatment(result)

## End(Not run)
```

requireDrugInDateRange

Restrict cohort to only cohort records within a certain date range

Description

Filter the cohort table keeping only the cohort records for which the specified index date is within a specified date range.

Usage

```
requireDrugInDateRange(
  cohort,
  dateRange,
  indexDate = "cohort_start_date",
  cohortId = NULL,
  name = omopgenerics::tableName(cohort)
)
```

Arguments

cohort	A cohort_table object.
dateRange	Date interval to consider. Any records with the index date outside of this range will be dropped.
indexDate	The column containing the date that will be checked against the date range.
cohortId	A cohort definition id to restrict by. If NULL, all cohorts will be included.
name	Name of the new cohort table, it must be a length 1 character vector.

Value

The cohort table having applied the date requirement.

Examples

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

cdm <- mockDrugUtilisation()

cdm$cohort1 <- cdm$cohort1 |>
  requireDrugInDateRange(dateRange = as.Date(c("2020-01-01", NA)))

attrition(cdm$cohort1) |>
  glimpse()
```

requireIsFirstDrugEntry

Restrict cohort to only the first cohort record per subject

Description

Filter the cohort table keeping only the first cohort record per subject.

Usage

```
requireIsFirstDrugEntry(
  cohort,
  cohortId = NULL,
  name = omopgenerics::tableName(cohort)
)
```

Arguments

cohort	A cohort_table object.
cohortId	A cohort definition id to restrict by. If NULL, all cohorts will be included.
name	Name of the new cohort table, it must be a length 1 character vector.

Value

The cohort table having applied the first entry requirement.

Examples

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

cdm <- mockDrugUtilisation()

cdm$cohort1 <- cdm$cohort1 |>
  requireIsFirstDrugEntry()

attrition(cdm$cohort1) |>
  glimpse()
```

requireObservationBeforeDrug

Restrict cohort to only cohort records with the given amount of prior observation time in the database

Description

Filter the cohort table keeping only the cohort records for which the individual has the required observation time in the database prior to their cohort start date.

Usage

```
requireObservationBeforeDrug(
  cohort,
  days,
  cohortId = NULL,
  name = omopgenerics::tableName(cohort)
)
```

Arguments

cohort	A cohort_table object.
days	Number of days of prior observation required before cohort start date. Any records with fewer days will be dropped.
cohortId	A cohort definition id to restrict by. If NULL, all cohorts will be included.
name	Name of the new cohort table, it must be a length 1 character vector.

Value

The cohort table having applied the prior observation requirement.

Examples

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

cdm <- mockDrugUtilisation()

cdm$cohort1 <- cdm$cohort1 |>
  requireObservationBeforeDrug(days = 365)

attrition(cdm$cohort1) |>
  glimpse()
```

requirePriorDrugWashout

Restrict cohort to only cohort records with a given amount of time since the last cohort record ended

Description

Filter the cohort table keeping only the cohort records for which the required amount of time has passed since the last cohort entry ended for that individual.

Usage

```
requirePriorDrugWashout(
  cohort,
  days,
  cohortId = NULL,
  name = omopgenerics::tableName(cohort)
)
```

Arguments

cohort	A cohort_table object.
days	The number of days required to have passed since the last cohort record finished. Any records with fewer days than this will be dropped. Note that setting days to Inf will lead to the same result as that from using the requireIsFirstDrugEntry function (with only an individual's first cohort record kept).
cohortId	A cohort definition id to restrict by. If NULL, all cohorts will be included.
name	Name of the new cohort table, it must be a length 1 character vector.

Value

The cohort table having applied the washout requirement.

Examples

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

cdm <- mockDrugUtilisation()

cdm$cohort1 <- cdm$cohort1 |>
  requirePriorDrugWashout(days = 90)

attrition(cdm$cohort1) |>
  glimpse()
```

summariseDoseCoverage *Check coverage of daily dose computation in a sample of the cdm for selected concept sets and ingredient*

Description

Check coverage of daily dose computation in a sample of the cdm for selected concept sets and ingredient

Usage

```
summariseDoseCoverage(
  cdm,
  ingredientConceptId,
  estimates = c("count_missing", "percentage_missing", "mean", "sd", "q25", "median",
    "q75"),
  sampleSize = NULL
)
```

Arguments

cdm	A <code>cdm_reference</code> object.
ingredientConceptId	Ingredient OMOP concept that we are interested for the study.
estimates	Estimates to obtain.
sampleSize	Maximum number of records of an ingredient to estimate dose coverage. If an ingredient has more, a random sample equal to <code>sampleSize</code> will be considered. If <code>NULL</code> , all records will be used.

Value

The function returns information of the coverage of `computeDailyDose.R` for the selected ingredients and concept sets

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation()

summariseDoseCoverage(cdm = cdm, ingredientConceptId = 1125315)
```

summariseDrugRestart *Summarise the drug restart for each follow-up period of interest.*

Description

Summarise the drug restart for each follow-up period of interest.

Usage

```
summariseDrugRestart(
  cohort,
  cohortId = NULL,
  switchCohortTable,
  switchCohortId = NULL,
  strata = list(),
  followUpDays = Inf,
  censorDate = NULL,
  incident = TRUE,
  restrictToFirstDiscontinuation = TRUE
)
```

Arguments

cohort	A cohort_table object.
cohortId	A cohort definition id to restrict by. If NULL, all cohorts will be included.
switchCohortTable	A cohort table in the cdm that contains possible alternative treatments.
switchCohortId	The cohort ids to be used from switchCohortTable. If NULL all cohort definition ids are used.
strata	A list of variables to stratify results. These variables must have been added as additional columns in the cohort table.
followUpDays	A vector of number of days to follow up. It can be multiple values.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
incident	Whether the switch treatment has to be incident (start after discontinuation) or not (it can start before the discontinuation and last till after).
restrictToFirstDiscontinuation	Whether to consider only the first discontinuation episode or all of them.

Value

A summarised_result object with the percentages of restart, switch and not exposed per follow-up period given.

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation()

conceptlist <- list(acetaminophen = 1125360, metformin = c(1503297, 1503327))
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "switch_cohort",
                                         conceptSet = conceptlist)

result <- cdm$cohort1 |>
  summariseDrugRestart(switchCohortTable = "switch_cohort")

tableDrugRestart(result)
```

summariseDrugUtilisation

This function is used to summarise the dose utilisation table over multiple cohorts.

Description

This function is used to summarise the dose utilisation table over multiple cohorts.

Usage

```
summariseDrugUtilisation(
  cohort,
  cohortId = NULL,
  strata = list(),
  estimates = c("q25", "median", "q75", "mean", "sd", "count_missing",
               "percentage_missing"),
  ingredientConceptId = NULL,
  conceptSet = NULL,
  indexDate = "cohort_start_date",
  censorDate = "cohort_end_date",
  restrictIncident = TRUE,
  gapEra = 1,
  numberExposures = TRUE,
  numberEras = TRUE,
  daysExposed = TRUE,
  daysPrescribed = TRUE,
```

```

    timeToExposure = TRUE,
    initialExposureDuration = TRUE,
    initialQuantity = TRUE,
    cumulativeQuantity = TRUE,
    initialDailyDose = TRUE,
    cumulativeDose = TRUE
)

```

Arguments

cohort	A cohort_table object.
cohortId	A cohort definition id to restrict by. If NULL, all cohorts will be included.
strata	A list of variables to stratify results. These variables must have been added as additional columns in the cohort table.
estimates	Estimates that we want for the columns.
ingredientConceptId	Ingredient OMOP concept that we are interested for the study.
conceptSet	List of concepts to be included.
indexDate	Name of a column that indicates the date to start the analysis.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
restrictIncident	Whether to include only incident prescriptions in the analysis. If FALSE all prescriptions that overlap with the study period will be included.
gapEra	Number of days between two continuous exposures to be considered in the same era.
numberExposures	Whether to include 'number_exposures' (number of drug exposure records between indexDate and censorDate).
numberEras	Whether to include 'number_eras' (number of continuous exposure episodes between indexDate and censorDate).
daysExposed	Whether to include 'days_exposed' (number of days that the individual is in a continuous exposure episode, including allowed treatment gaps, between indexDate and censorDate; sum of the length of the different drug eras).
daysPrescribed	Whether to include 'days_prescribed' (sum of the number of days for each prescription that contribute in the analysis).
timeToExposure	Whether to include 'time_to_exposure' (number of days between indexDate and the first episode).
initialExposureDuration	Whether to include 'initial_exposure_duration' (number of prescribed days of the first drug exposure record).
initialQuantity	Whether to include 'initial_quantity' (quantity of the first drug exposure record).
cumulativeQuantity	Whether to include 'cumulative_quantity' (sum of the quantity of the different exposures considered in the analysis).

initialDailyDose
 Whether to include 'initial_daily_dose_{unit}' (daily dose of the first considered prescription).

cumulativeDose Whether to include 'cumulative_dose_{unit}' (sum of the cumulative dose of the analysed drug exposure records).

Value

A summary of drug utilisation stratified by cohort_name and strata_name

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)

cdm <- mockDrugUtilisation()
cdm <- generateIngredientCohortSet(cdm = cdm,
                                      ingredient = "acetaminophen",
                                      name = "dus_cohort")

cdm$dus_cohort |>
  summariseDrugUtilisation(ingredientConceptId = 1125315)
```

summariseIndication *Summarise the indications of individuals in a drug cohort*

Description

Summarise the observed indications of patients in a drug cohort based on their presence in an indication cohort in a specified time window. If an individual is not in one of the indication cohorts, they will be considered to have an unknown indication if they are present in one of the specified OMOP CDM clinical tables. Otherwise, if they are neither in an indication cohort or a clinical table they will be considered as having no observed indication.

Usage

```
summariseIndication(
  cohort,
  strata = list(),
  indicationCohortName,
  cohortId = NULL,
  indicationCohortId = NULL,
  indicationWindow = list(c(0, 0)),
  unknownIndicationTable = NULL,
  indexDate = "cohort_start_date",
  mutuallyExclusive = TRUE,
  censorDate = NULL
)
```

Arguments

cohort	A cohort_table object.
strata	A list of variables to stratify results. These variables must have been added as additional columns in the cohort table.
indicationCohortName	Name of the cohort table with potential indications.
cohortId	A cohort definition id to restrict by. If NULL, all cohorts will be included.
indicationCohortId	The target cohort ID to add indication. If NULL all cohorts will be considered.
indicationWindow	The time window over which to identify indications.
unknownIndicationTable	Tables in the OMOP CDM to search for unknown indications.
indexDate	Name of a column that indicates the date to start the analysis.
mutuallyExclusive	Whether to report indications as mutually exclusive or report them as independent results.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.

Value

A summarised result

Examples

```
library(DrugUtilisation)
library(CDMConnector)
library(dplyr, warn.conflicts = FALSE)

cdm <- mockDrugUtilisation()

indications <- list(headache = 378253, asthma = 317009)
cdm <- generateConceptCohortSet(cdm = cdm,
                                   conceptSet = indications,
                                   name = "indication_cohorts")

cdm <- generateIngredientCohortSet(cdm = cdm,
                                       name = "drug_cohort",
                                       ingredient = "acetaminophen")

cdm$drug_cohort |>
  summariseIndication(
    indicationCohortName = "indication_cohorts",
    unknownIndicationTable = "condition_occurrence",
    indicationWindow = list(c(-Inf, 0))
  ) |>
  glimpse()
```

summariseProportionOfPatientsCovered
Summarise proportion Of patients covered

Description

Gives the proportion of patients still in observation who are in the cohort on any given day following their first cohort entry. This is known as the “proportion of patients covered” (PPC) method for assessing treatment persistence.

Usage

```
summariseProportionOfPatientsCovered(
  cohort,
  cohortId = NULL,
  strata = list(),
  followUpDays = NULL
)
```

Arguments

<code>cohort</code>	A cohort_table object.
<code>cohortId</code>	A cohort definition id to restrict by. If NULL, all cohorts will be included.
<code>strata</code>	A list of variables to stratify results. These variables must have been added as additional columns in the cohort table.
<code>followUpDays</code>	Number of days to follow up individuals for. If NULL the maximum amount of days from an individuals first cohort start date to their last cohort end date will be used

Value

A summarised result

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation(numberIndividuals = 100)

result <- cdm$cohort1 |>
  summariseProportionOfPatientsCovered(followUpDays = 365)

tidy(result)
```

summariseTreatment *This function is used to summarise treatments received*

Description

This function is used to summarise treatments received

Usage

```
summariseTreatment(  
  cohort,  
  window,  
  treatmentCohortName,  
  cohortId = NULL,  
  treatmentCohortId = NULL,  
  strata = list(),  
  indexDate = "cohort_start_date",  
  censorDate = NULL,  
  mutuallyExclusive = FALSE  
)
```

Arguments

cohort	A cohort_table object.
window	Time window over which to summarise the treatments.
treatmentCohortName	Name of a cohort in the cdm that contains the treatments of interest.
cohortId	A cohort definition id to restrict by. If NULL, all cohorts will be included.
treatmentCohortId	Cohort definition id of interest from treatmentCohortName.
strata	A list of variables to stratify results. These variables must have been added as additional columns in the cohort table.
indexDate	Name of a column that indicates the date to start the analysis.
censorDate	Name of a column that indicates the date to stop the analysis, if NULL end of individuals observation is used.
mutuallyExclusive	Whether to include mutually exclusive treatments or not.

Value

A summary of treatments stratified by cohort_name and strata_name

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation()
cdm$cohort1 |>
  summariseTreatment(
    treatmentCohortName = "cohort2",
    window = list(c(0, 30), c(31, 365))
)
```

tableDoseCoverage *Format a dose_coverage object into a visual table.*

Description

Format a dose_coverage object into a visual table.

Usage

```
tableDoseCoverage(
  result,
  header = c("variable_name", "estimate_name"),
  groupColumn = c("cdm_name", "ingredient_name"),
  type = "gt",
  hide = c("variable_level", "sample_size"),
  .options = list()
)
```

Arguments

<code>result</code>	A summarised_result object.
<code>header</code>	Columns to use as header. See options with availableTableColumns(result).
<code>groupColumn</code>	Columns to group by. See options with availableTableColumns(result).
<code>type</code>	Type of table. Check supported types with visOmopResults::tableType().
<code>hide</code>	Columns to hide from the visualisation. See options with availableTableColumns(result).
<code>.options</code>	A named list with additional formatting options. visOmopResults::tableOptions() shows allowed arguments and their default values.

Value

A table with a formatted version of summariseDrugCoverage() results.

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation()

result <- summariseDoseCoverage(cdm, 1125315)

tableDoseCoverage(result)
```

tableDrugRestart *Format a drug_restart object into a visual table.*

Description

Format a drug_restart object into a visual table.

Usage

```
tableDrugRestart(
  result,
  header = c("cdm_name", "cohort_name"),
  groupColumn = "variable_name",
  type = "gt",
  hide = c("censor_date", "restrict_to_first_discontinuation", "follow_up_days",
    "cohort_table_name", "incident", "switch_cohort_table"),
  .options = list()
)
```

Arguments

result	A summarised_result object.
header	Columns to use as header. See options with availableTableColumns(result).
groupColumn	Columns to group by. See options with availableTableColumns(result).
type	Type of table. Check supported types with visOmopResults::tableType().
hide	Columns to hide from the visualisation. See options with availableTableColumns(result).
.options	A named list with additional formatting options. visOmopResults::tableOptions() shows allowed arguments and their default values.

Value

A table with a formatted version of summariseDrugRestart() results.

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation()

conceptlist <- list(acetaminophen = 1125360, metformin = c(1503297, 1503327))
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "switch_cohort",
                                         conceptSet = conceptlist)

result <- cdm$cohort1 |>
  summariseDrugRestart(switchCohortTable = "switch_cohort")

tableDrugRestart(result)
```

tableDrugUtilisation *Format a drug_utilisation object into a visual table.*

Description

Format a drug_utilisation object into a visual table.

Usage

```
tableDrugUtilisation(
  result,
  header = c("cdm_name"),
  groupColumn = c("cohort_name", strataColumns(result)),
  type = "gt",
  hide = c("variable_level", "censor_date", "cohort_table_name", "gap_era", "index_date",
          "restrict_incident"),
  .options = list()
)
```

Arguments

- result** A summarised_result object.
- header** Columns to use as header. See options with availableTableColumns(result).
- groupColumn** Columns to group by. See options with availableTableColumns(result).
- type** Type of table. Check supported types with visOmopResults::tableType().
- hide** Columns to hide from the visualisation. See options with availableTableColumns(result).
- .options** A named list with additional formatting options. visOmopResults::tableOptions() shows allowed arguments and their default values.

Value

A table with a formatted version of summariseIndication() results.

Examples

```
library(DrugUtilisation)
library(CodelistGenerator)

cdm <- mockDrugUtilisation()
codelist <- getDrugIngredientCodes(cdm = cdm, name = "acetaminophen")
cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "dus_cohort",
                                         conceptSet = codelist)

drugUse <- cdm$dus_cohort |>
  summariseDrugUtilisation(ingredientConceptId = 1125315)

tableDrugUtilisation(drugUse)
```

tableIndication	<i>Create a table showing indication results</i>
-----------------	--

Description

Create a table showing indication results

Usage

```
tableIndication(
  result,
  header = c("cdm_name", "cohort_name", strataColumns(result)),
  groupColumn = "variable_name",
  hide = c("window_name", "mutually_exclusive", "unknown_indication_table",
           "censor_date", "cohort_table_name", "index_date", "indication_cohort_name"),
  type = "gt",
  .options = list()
)
```

Arguments

<code>result</code>	A summarised_result object.
<code>header</code>	Columns to use as header. See options with availableTableColumns(result).
<code>groupColumn</code>	Columns to group by. See options with availableTableColumns(result).
<code>hide</code>	Columns to hide from the visualisation. See options with availableTableColumns(result).
<code>type</code>	Type of table. Check supported types with visOmopResults::tableType().
<code>.options</code>	A named list with additional formatting options. visOmopResults::tableOptions() shows allowed arguments and their default values.

Value

A table with a formatted version of summariseIndication() results.

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation()

result <- cdm$cohort1 |>
  summariseIndication(
    indicationCohortName = "cohort2",
    indicationWindow = list(c(-30, 0)),
    unknownIndicationTable = "condition_occurrence"
  )

tableIndication(result)
```

tableProportionOfPatientsCovered

Create a table with proportion of patients covered results

Description

Create a table with proportion of patients covered results

Usage

```
tableProportionOfPatientsCovered(
  result,
  header = c("cohort_name", strataColumns(result)),
  groupColumn = "cdm_name",
  type = "gt",
  hide = c("variable_name", "variable_level", "cohort_table_name"),
  .options = list()
)
```

Arguments

- | | |
|--------------------------|---|
| <code>result</code> | A summarised_result object. |
| <code>header</code> | Columns to use as header. See options with availableTableColumns(result). |
| <code>groupColumn</code> | Columns to group by. See options with availableTableColumns(result). |
| <code>type</code> | Type of table. Check supported types with visOmopResults::tableType(). |
| <code>hide</code> | Columns to hide from the visualisation. See options with availableTableColumns(result). |
| <code>.options</code> | A named list with additional formatting options. visOmopResults::tableOptions() shows allowed arguments and their default values. |

Value

A table with a formatted version of summariseProportionOfPatientsCovered() results.

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation()

cdm <- generateDrugUtilisationCohortSet(cdm = cdm,
                                         name = "my_cohort",
                                         conceptSet = list(drug_of_interest = c(1503297, 1503327)))

result <- cdm$my_cohort |>
  summariseProportionOfPatientsCovered(followUpDays = 365)

tableProportionOfPatientsCovered(result)
```

tableTreatment*Format a summarised_treatment result into a visual table.*

Description

Format a summarised_treatment result into a visual table.

Usage

```
tableTreatment(
  result,
  header = c("cdm_name", "cohort_name"),
  groupColumn = "variable_name",
  type = "gt",
  hide = c("window_name", "mutually_exclusive", "censor_date", "cohort_table_name",
          "index_date", "treatment_cohort_name"),
  .options = list()
)
```

Arguments

<code>result</code>	A summarised_result object.
<code>header</code>	Columns to use as header. See options with availableTableColumns(result).
<code>groupColumn</code>	Columns to group by. See options with availableTableColumns(result).
<code>type</code>	Type of table. Check supported types with visOmopResults::tableType().
<code>hide</code>	Columns to hide from the visualisation. See options with availableTableColumns(result).
<code>.options</code>	A named list with additional formatting options. visOmopResults::tableOptions() shows allowed arguments and their default values.

Value

A table with a formatted version of summariseTreatment() results.

Examples

```
library(DrugUtilisation)

cdm <- mockDrugUtilisation()

result <- cdm$cohort1 |>
  summariseTreatment(
    treatmentCohortName = "cohort2",
    window = list(c(0, 30), c(31, 365))
  )

tableTreatment(result)
```

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