

# Package ‘OmopSketch’

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**Title** Characterise Tables of an OMOP Common Data Model Instance

**Version** 0.1.2

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**Description** Summarises key information in data mapped to the Observational Medical Outcomes Partnership (OMOP) common data model. Assess suitability to perform specific epidemiological studies and explore the different domains to obtain feasibility counts and trends.

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

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**Depends** R (>= 2.10)

**URL** <https://OHDSI.github.io/OmopSketch/>

**BugReports** <https://github.com/OHDSI/OmopSketch/issues>

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**mockOmopSketch**      *Creates a mock database to test OmopSketch package.*

### Description

Creates a mock database to test OmopSketch package.

### Usage

```
mockOmopSketch(
  con = NULL,
  writeSchema = NULL,
  numberIndividuals = 100,
  seed = NULL
)
```

### Arguments

<code>con</code>	A DBI connection to create the cdm mock object. By default, the connection would be a 'duckdb' one.
<code>writeSchema</code>	Name of an schema of the DBI connection with writing permissions.
<code>numberIndividuals</code>	Number of individuals to create in the cdm reference object.
<code>seed</code>	An optional integer used to set the seed for random number generation, ensuring reproducibility of the generated data. If provided, this seed allows the function to produce consistent results each time it is run with the same inputs. If 'NULL', the seed is not set, which can lead to different outputs on each run.

**Value**

A mock cdm\_reference object.

**Examples**

```
library(OmopSketch)
mockOmopSketch(numberIndividuals = 100)
```

---

plotConceptCounts

*Plot the concept counts of a summariseConceptCounts output.*

---

**Description**

Plot the concept counts of a summariseConceptCounts output.

**Usage**

```
plotConceptCounts(result, facet = NULL, colour = NULL)
```

**Arguments**

- |        |  |
|--------|--|
| result | A summarised_result object (output of summariseConceptCounts).   |
| facet  | Columns to face by. Formula format can be provided. See possible columns to face by with: visOmopResults::tidyColumns(). |
| colour | Columns to colour by. See possible columns to colour by with: visOmopResults::tidyColumns().                             |

**Value**

A ggplot2 object showing the concept counts.

**Examples**

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

cdm <- mockOmopSketch()

result <- cdm |>
  summariseConceptCounts(
    conceptId = list(
      "Renal agenesis" = 194152,
      "Manic mood" = c(4226696, 4304866, 37110496, 40371897)
    )
  )

result |>
  filter(estimate_name == "person_count", variable_name == "overall") |>
  plotConceptCounts(facet = "codelist_name", colour = "codelist_name")
```

```
PatientProfiles::mockDisconnect(cdm)
```

**plotInObservation**      *Create a ggplot2 plot from the output of summariseInObservation().*

## Description

Create a ggplot2 plot from the output of summariseInObservation().

## Usage

```
plotInObservation(result, facet = NULL, colour = NULL)
```

## Arguments

- result**      A summarised\_result object (output of summariseInObservation).
- facet**      Columns to face by. Formula format can be provided. See possible columns to face by with: visOmopResults::tidyColumns().
- colour**      Columns to colour by. See possible columns to colour by with: visOmopResults::tidyColumns().

## Value

A ggplot showing the table counts

## Examples

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

cdm <- mockOmopSketch()

result <- summariseInObservation(
  cdm$observation_period,
  output = c("person-days", "records"),
  ageGroup = list("<=40" = c(0, 40), ">40" = c(41, Inf)),
  sex = TRUE
)

result |>
  filter(variable_name == "Number person-days") |>
  plotInObservation(facet = "sex", colour = "age_group")

PatientProfiles::mockDisconnect(cdm)
```

---

plotObservationPeriod *Create a plot from the output of summariseObservationPeriod().*

---

## Description

Create a plot from the output of summariseObservationPeriod().

## Usage

```
plotObservationPeriod(  
  result,  
  variableName = "number subjects",  
  plotType = "barplot",  
  facet = NULL,  
  colour = NULL  
)
```

## Arguments

result	A summarised_result object.
variableName	The variable to plot it can be: "number subjects", "records per person", "duration" or "days to next observation period".
plotType	The plot type, it can be: "barplot", "boxplot" or "densityplot".
facet	Columns to colour by. See possible columns to colour by with: visOmopResults::tidyColumns().
colour	Columns to colour by. See possible columns to colour by with: visOmopResults::tidyColumns().

## Value

A ggplot2 object.

## Examples

```
cdm <- mockOmopSketch(numberIndividuals = 100)  
  
result <- summariseObservationPeriod(cdm$observation_period)  
  
result |>  
  plotObservationPeriod(  
    variableName = "duration in days",  
    plotType = "boxplot"  
)  
  
PatientProfiles::mockDisconnect(cdm)
```

`plotRecordCount`      *Create a ggplot of the records' count trend.*

## Description

Create a ggplot of the records' count trend.

## Usage

```
plotRecordCount(result, facet = NULL, colour = NULL)
```

## Arguments

<code>result</code>	Output from <code>summariseRecordCount()</code> .
<code>facet</code>	Columns to face by. Formula format can be provided. See possible columns to face by with: <code>visOmopResults::tidyColumns()</code> .
<code>colour</code>	Columns to colour by. See possible columns to colour by with: <code>visOmopResults::tidyColumns()</code> .

## Value

A ggplot showing the table counts

## Examples

```
cdm <- mockOmopSketch()

summarisedResult <- summariseRecordCount(
  cdm = cdm,
  omopTableName = "condition_occurrence",
  ageGroup = list("<=20" = c(0,20), ">20" = c(21, Inf)),
  sex = TRUE
)

plotRecordCount(summarisedResult, colour = "age_group", facet = sex ~ .)

PatientProfiles::mockDisconnect(cdm = cdm)
```

## summariseClinicalRecords

*Summarise an omop table from a cdm object. You will obtain information related to the number of records, number of subjects, whether the records are in observation, number of present domains and number of present concepts.*

## Description

Summarise an omop table from a cdm object. You will obtain information related to the number of records, number of subjects, whether the records are in observation, number of present domains and number of present concepts.

## Usage

```
summariseClinicalRecords(
  cdm,
  omopTableName,
  recordsPerPerson = c("mean", "sd", "median", "q25", "q75", "min", "max"),
  inObservation = TRUE,
  standardConcept = TRUE,
  sourceVocabulary = FALSE,
  domainId = TRUE,
  typeConcept = TRUE,
  sex = FALSE,
  ageGroup = NULL
)
```

## Arguments

<code>cdm</code>	A <code>cdm_reference</code> object.
<code>omopTableName</code>	A character vector of the names of the tables to summarise in the <code>cdm</code> object.
<code>recordsPerPerson</code>	Generates summary statistics for the number of records per person. Set to <code>NULL</code> if no summary statistics are required.
<code>inObservation</code>	Boolean variable. Whether to include the percentage of records in observation.
<code>standardConcept</code>	Boolean variable. Whether to summarise standard concept information.
<code>sourceVocabulary</code>	Boolean variable. Whether to summarise source vocabulary information.
<code>domainId</code>	Boolean variable. Whether to summarise domain id of standard concept id information.
<code>typeConcept</code>	Boolean variable. Whether to summarise type concept id field information.
<code>sex</code>	Boolean variable. Whether to stratify by sex (TRUE) or not (FALSE).
<code>ageGroup</code>	A list of age groups to stratify results by.

## Value

A `summarised_result` object.

## Examples

```
cdm <- mockOmopSketch()

summarisedResult <- summariseClinicalRecords(
```

```

cdm = cdm,
omopTableName = "condition_occurrence",
recordsPerPerson = c("mean", "sd"),
inObservation = TRUE,
standardConcept = TRUE,
sourceVocabulary = TRUE,
domainId = TRUE,
typeConcept = TRUE
)
summarisedResult
PatientProfiles::mockDisconnect(cdm = cdm)

```

**summariseConceptCounts***Summarise code use in patient-level data***Description**

Summarise code use in patient-level data

**Usage**

```

summariseConceptCounts(
  cdm,
  conceptId,
  countBy = c("record", "person"),
  concept = TRUE,
  year = FALSE,
  sex = FALSE,
  ageGroup = NULL
)

```

**Arguments**

<code>cdm</code>	A <code>cdm</code> object
<code>conceptId</code>	List of concept IDs to summarise.
<code>countBy</code>	Either "record" for record-level counts or "person" for person-level counts
<code>concept</code>	TRUE or FALSE. If TRUE code use will be summarised by concept.
<code>year</code>	TRUE or FALSE. If TRUE code use will be summarised by year.
<code>sex</code>	TRUE or FALSE. If TRUE code use will be summarised by sex.
<code>ageGroup</code>	A list of <code>ageGroup</code> vectors of length two. Code use will be thus summarised by age groups.

**Value**

A summarised\_result object with results overall and, if specified, by strata.

**Examples**

```
cdm <- mockOmopSketch()

cs <- list(sumatriptan = c(35604883, 35604879, 35604880, 35604884))

results <- summariseConceptCounts(cdm, conceptId = cs)

results

PatientProfiles::mockDisconnect(cdm)
```

**summariseInObservation**

*Summarise the number of people in observation during a specific interval of time.*

**Description**

Summarise the number of people in observation during a specific interval of time.

**Usage**

```
summariseInObservation(
  observationPeriod,
  unit = "year",
  unitInterval = 1,
  output = "records",
  ageGroup = NULL,
  sex = FALSE
)
```

**Arguments**

observationPeriod	An observation_period omop table. It must be part of a cdm_reference object.
unit	Whether to stratify by "year" or by "month".
unitInterval	Number of years or months to include within the time interval.
output	Output format. It can be either the number of records ("records") that are in observation in the specific interval of time, the number of person-days ("person-days"), or both c("records","person-days").
ageGroup	A list of age groups to stratify results by.
sex	Boolean variable. Whether to stratify by sex (TRUE) or not (FALSE).

**Value**

A summarised\_result object.

**Examples**

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

cdm <- mockOmopSketch()

result <- summariseInObservation(
  cdm$observation_period,
  unit = "month",
  unitInterval = 6,
  output = c("person-days", "records"),
  ageGroup = list("<=60" = c(0, 60), ">60" = c(61, Inf)),
  sex = TRUE
)

result |>
  glimpse()

PatientProfiles::mockDisconnect(cdm)
```

**summariseObservationPeriod**

*Summarise the observation period table getting some overall statistics in a summarised\_result object.*

**Description**

Summarise the observation period table getting some overall statistics in a summarised\_result object.

**Usage**

```
summariseObservationPeriod(
  observationPeriod,
  estimates = c("mean", "sd", "min", "q05", "q25", "median", "q75", "q95", "max",
    "density"),
  ageGroup = NULL,
  sex = FALSE
)
```

**Arguments**

**observationPeriod**  
observation\_period omop table.

estimates	Estimates to summarise the variables of interest ( records per person, duration in days and days to next observation period).
ageGroup	A list of age groups to stratify results by.
sex	Boolean variable. Whether to stratify by sex (TRUE) or not (FALSE).

**Value**

A summarised\_result object with the summarised data.

**Examples**

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

cdm <- mockOmopSketch(numberIndividuals = 100)

result <- summariseObservationPeriod(cdm$observation_period)

result |>
  glimpse()

PatientProfiles::mockDisconnect(cdm)
```

**summariseOmopSnapshot** *Summarise a cdm\_reference object creating a snapshot with the metadata of the cdm\_reference object.*

**Description**

Summarise a cdm\_reference object creating a snapshot with the metadata of the cdm\_reference object.

**Usage**

```
summariseOmopSnapshot(cdm)
```

**Arguments**

cdm	A cdm_reference object.
-----	-------------------------

**Value**

A summarised\_result object.

## Examples

```
library(OmopSketch)
cdm <- mockOmopSketch(numberIndividuals = 10)

summariseOmopSnapshot(cdm)
```

**summarisePopulationCharacteristics**

*Summarise the characteristics of the base population of a cdm\_reference object.*

## Description

Summarise the characteristics of the base population of a cdm\_reference object.

## Usage

```
summarisePopulationCharacteristics(
  cdm,
  studyPeriod = c(NA, NA),
  sex = FALSE,
  ageGroup = NULL
)
```

## Arguments

cdm	A cdm_reference object.
studyPeriod	Dates to trim the observation period. If NA, min(observation_period_start_date) and/or max(observation_period_end_date) are used.
sex	Whether to stratify the results by sex.
ageGroup	List of age groups to stratify by at index date.

## Value

A summarised\_result object.

## Examples

```
cdm <- mockOmopSketch()

summarisedPopulation <- summarisePopulationCharacteristics(
  cdm = cdm,
  studyPeriod = c("2010-01-01", NA),
  sex = TRUE,
  ageGroup = NULL
)
```

```
summarisedPopulation |> print()

PatientProfiles::mockDisconnect(cdm = cdm)
```

**summariseRecordCount** *Summarise record counts of an omop\_table using a specific time interval. Only records that fall within the observation period are considered.*

## Description

Summarise record counts of an omop\_table using a specific time interval. Only records that fall within the observation period are considered.

## Usage

```
summariseRecordCount(
  cdm,
  omopTableName,
  unit = "year",
  unitInterval = 1,
  ageGroup = NULL,
  sex = FALSE
)
```

## Arguments

cdm	A cdm_reference object.
omopTableName	A character vector of omop tables from the cdm.
unit	Time unit it can either be "year" or "month".
unitInterval	Number of years or months to include within the same interval.
ageGroup	A list of age groups to stratify results by.
sex	Whether to stratify by sex (TRUE) or not (FALSE).

## Value

A summarised\_result object.

## Examples

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

cdm <- mockOmopSketch()

summarisedResult <- summariseRecordCount(
```

```

cdm = cdm,
omopTableName = c("condition_occurrence", "drug_exposure"),
unit = "year",
unitInterval = 10,
ageGroup = list("<=20" = c(0,20), ">20" = c(21, Inf)),
sex = TRUE
)

summarisedResult |>
glimpse()

PatientProfiles::mockDisconnect(cdm = cdm)

```

**tableClinicalRecords** *Create a visual table from a summariseClinicalRecord() output.*

## Description

Create a visual table from a summariseClinicalRecord() output.

## Usage

```
tableClinicalRecords(result, type = "gt")
```

## Arguments

result	Output from summariseClinicalRecords().
type	Type of formatting output table, either "gt" or "flextable".

## Value

A gt or flextable object with the summarised data.

## Examples

```

cdm <- mockOmopSketch()

summarisedResult <- summariseClinicalRecords(
  cdm = cdm,
  omopTableName = c("condition_occurrence", "drug_exposure"),
  recordsPerPerson = c("mean", "sd"),
  inObservation = TRUE,
  standardConcept = TRUE,
  sourceVocabulary = TRUE,
  domainId = TRUE,
  typeConcept = TRUE
)

```

```
summarisedResult |>
  suppress(minCellCount = 5) |>
  tableClinicalRecords()

PatientProfiles::mockDisconnect(cdm)
```

---

**tableObservationPeriod**

*Create a visual table from a summariseObservationPeriod() result.*

---

**Description**

Create a visual table from a summariseObservationPeriod() result.

**Usage**

```
tableObservationPeriod(result, type = "gt")
```

**Arguments**

result	A summarised_result object.
type	Type of formatting output table, either "gt" or "flextable".

**Value**

A gt or flextable object with the summarised data.

**Examples**

```
cdm <- mockOmopSketch(numberIndividuals = 100)

result <- summariseObservationPeriod(cdm$observation_period)

tableObservationPeriod(result)

PatientProfiles::mockDisconnect(cdm)
```

**tableOmopSnapshot**      *Create a visual table from a summarise\_omop\_snapshot result.*

## Description

Create a visual table from a summarise\_omop\_snapshot result.

## Usage

```
tableOmopSnapshot(result, type = "gt")
```

## Arguments

result	Output from summariseOmopSnapshot().
type	Type of formatting output table, either "gt" or "flextable".

## Value

A gt or flextable object with the summarised data.

## Examples

```
library(OmopSketch)
cdm <- mockOmopSketch(numberIndividuals = 10)

result <- summariseOmopSnapshot(cdm)

result |>
  tableOmopSnapshot()

PatientProfiles::mockDisconnect(cdm)
```

**tablePopulationCharacteristics**

*Create a visual table from a summarise\_population\_characteristics result.*

## Description

Create a visual table from a summarise\_population\_characteristics result.

## Usage

```
tablePopulationCharacteristics(result, type = "gt")
```

**Arguments**

result	Output from summarisePopulationCharacteristics().
type	Type of formatting output table, either "gt" or "flextable".

**Value**

A gt or flextable object with the summarised data.

**Examples**

```
cdm <- mockOmopSketch()

summarisedPopulation <- summarisePopulationCharacteristics(
  cdm = cdm,
  studyPeriod = c("2010-01-01", NA),
  sex = TRUE,
  ageGroup = list("<=60" = c(0, 60), ">60" = c(61, Inf))
)

summarisedPopulation |>
  suppress(minCellCount = 5) |>
  tablePopulationCharacteristics()

PatientProfiles::mockDisconnect(cdm = cdm)
```

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