# Package 'PhenotypeLibrary'

September 23, 2023

Type Package
Title The OHDSI Phenotype Library
Version 3.22.0
<b>Date</b> 2023-09-23
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<b>Description</b> A repository to store the content of the OHDSI Phenotype library.
Imports checkmate, readr, dplyr, rlang
Suggests reactable, testthat, knitr
License Apache License
RoxygenNote 7.2.3
Roxygen list(markdown = TRUE)
Encoding UTF-8
Language en-US
VignetteBuilder knitr
<pre>URL https://ohdsi.github.io/PhenotypeLibrary/, https:    //github.com/OHDSI/PhenotypeLibrary</pre>
BugReports https://github.com/OHDSI/PhenotypeLibrary/issues
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 ${\tt getPhenotypeLog}$ 

Get phenotype log

# Description

Get phenotype log

#### Usage

```
getPhenotypeLog(cohortIds = listPhenotypes()$cohortId)
```

# Arguments

cohortIds

IDs of cohorts to extraction from the library.

#### Value

Returns a table with one row per cohort definitions with log information such as its release cycle. Example, this function gives us insight on when a cohort definition was added/updated/deprecated by the OHDSI PhenotypeLibrary.

A tibble.

# **Examples**

```
getPhenotypeLog(cohortIds = c(1, 2))
```

```
getPlCohortDefinitionSet
```

Get a cohort definition set

# Description

Get a cohort definition set

#### Usage

```
getPlCohortDefinitionSet(cohortIds)
```

#### **Arguments**

cohortIds

IDs of cohorts to extraction from the library.

# Value

A tibble with the cohort ID, name, sql, and JSON for the provided cohort IDs. Can be used by the CohortGenerator package.

#### **Examples**

```
cohorts <- listPhenotypes()
subsetIds <- cohorts$cohortId[1:3]
getPlCohortDefinitionSet(subsetIds)</pre>
```

```
{\tt getPlConceptDefinitionSet}
```

Get conceptSets in cohorts

# Description

Get conceptSets in cohorts

# Usage

```
getPlConceptDefinitionSet(cohortIds = listPhenotypes()$cohortId)
```

# Arguments

cohortIds

IDs of cohorts to extraction from the library.

#### Value

Returns a table with one row per concept set for given cohort definitions.

A tibble.

# Examples

```
getPhenotypeLog(cohortIds = c(1, 2))
```

listPhenotypes

List all phenotypes in the library

# Description

List all phenotypes in the library

#### Usage

listPhenotypes()

#### Value

A tibble with the cohort ID and name.

# Examples

```
listPhenotypes()
```

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