# How to Use PhenotypeLibrary R Package

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| Ph | nenotypeLibrary is part of HADES |   |

#### 1 Installation

• This is an installable R-package that may be installed as follows:

```
remotes::install_github("OHDSI/PhenotypeLibrary")
```

#### 2 Retrieval

• The list of cohort definitions available may be retrieved as follows:

```
PhenotypeLibrary::getPhenotypeLog()
```

```
#> # A tibble: 563 x 81
#>
      cohortId cohortName
                           cohortNameAtlas cohortNameFormatted cohortNameLong librarian status addedVer
#>
         <dbl> <chr>
                           <chr>
                                                                <chr>
                                                                               <chr>>
                                                                                          <chr>
             3 Cough or S~ [P] Cough or S~ Cough or Sputum
#>
   1
                                                                Cough or Sput~ rao@ohds~ Pendi~ <NA>
#>
   2
             4 Diarrhea
                           [P] Diarrhea
                                           Diarrhea
                                                                Diarrhea
                                                                               rao@ohds~ Pendi~ <NA>
             5 Dyspnea
                           [P] Dyspnea
                                                                               rao@ohds~ Pendi~ <NA>
#>
   3
                                           Dyspnea
                                                                Dyspnea
#>
   4
             6 Fever
                           [P] Fever
                                           Fever
                                                                Fever
                                                                               rao@ohds~ Pendi~ <NA>
#>
             7 Headache o~ [P] Headache o~ Headache or Headache or H~ rao@ohds~ Pendi~ <NA>
#>
   6
             8 Altered sm~ [P] Altered sm~ Altered smell or t~ Altered smell~ rao@ohds~ Pendi~ <NA>
             9 Sore throat [P] Sore throat Sore throat
                                                                Sore throat
                                                                               rao@ohds~ Pendi~ <NA>
            10 Nausea or ~ [P] Nausea or ~ Nausea or Vomiting Nausea or Vom~ rao@ohds~ Pendi~ <NA>
            11 Malaise an~ [P] Malaise an~ Malaise and or fat~ Malaise and o~ rao@ohds~ Pendi~ <NA>
#> 10
            12 Rhinitis o~ [P] Rhinitis o~ Rhinitis or common~ Rhinitis or c~ rao@ohds~ Pendi~ <NA>
#> # i 553 more rows
```

```
#> # i 72 more variables: hashTag <chr>, isCirceJson <dbl>, contributors <chr>, contributorOrcIds <chr>
#> # contributorOrganizations <chr>, peerReviewers <chr>, peerReviewerOrcIds <dbl>, recommendedReferent
#> # ohdsiForumPost <chr>, createdDate <date>, modifiedDate <date>, lastModifiedBy <dbl>, replaces <dinterplaces <dinterplace
```

• You can extract one or more cohort definitions into a cohortDefinitionSet object as

```
cohortDefinitionSet <- PhenotypeLibrary::getPlCohortDefinitionSet(cohortIds = c(1, 2, 3))
cohortDefinitionSet</pre>
```

• cohortDefinitionSet is now a data.frame with specifications for the cohort ids 1, 2 and 3. For cohorts that conform to OHDSI Circe specifications, the field json is the cohort json specification that may be posted into your Atlas instance. The SQL is the SQL rendered from the JSON. For cohorts that do not conform to OHDSI Circe specification, only the SQL is provided and the json is left empty.

## 3 Use

• You can instantiate the cohorts in your environment as follows using (OHDSI/CohortGenerator)[https://github.com/OHDSI/CohortGenerator].

```
connectionDetails <-
  DatabaseConnector::createConnectionDetails(
  dbms = "postgresq1",
  server = "some.server.com/ohdsi",
  user = "joe",
  password = "secret"
)

cdmDatabaseSchema <- "cdm_synpuf"
cohortDatabaseSchema <- "scratch.dbo"
cohortTables <- CohortGenerator::getCohortTableNames()

CohortGenerator::generateCohortSet(
  connectionDetails = connectionDetails,
  cdmDatabaseSchema = cdmDatabaseSchema,
  cohortDatabaseSchema = cohortDatabaseSchema,
  cohortTableNames = cohortTables,
  cohortDefinitionSet = cohortDefinitionSet
)</pre>
```

• You can also run cohort diagnostics on this cohortDefinitionSet object as follows:

```
databaseId <- "synpuf"

databaseName <-
    "Medicare Claims Synthetic Public Use Files (SynPUFs)"

databaseDescription <-
    "Medicare Claims Synthetic Public Use Files (SynPUFs) were created to allow interested parties to gain

CohortDiagnostics::executeDiagnostics(
    cohortDefinitionSet = cohortDefinitionSet,
    exportFolder = outputFolder,
    databaseId = databaseId,
    databaseDescription = databaseDescription,
    cohortDatabaseSchema = cohortDatabaseSchema,
    codmDatabaseSchema = cohortDatabaseSchema,
    connectionDetails = connectionDetails,
    cohortTableNames = cohortTableNames
)</pre>
```