

# Package ‘OmicsMLRepoR’

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**Type** Package

**Title** Search harmonized metadata created under the OmicsMLRepo project

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**Description** This package provides functions to browse the harmonized metadata for large omics databases. This package also supports data navigation if the metadata incorporates ontology.

**Depends** R (>= 4.4.0)

**Imports** dplyr, stringr, rols, tidyr, methods, stats, tibble, data.tree, jsonlite, plyr, BiocFileCache, readr, DiagrammeR, rlang, lubridate

**Suggests** arrow, knitr, BiocStyle, curatedMetagenomicData, testthat (>= 3.0.0), cBioPortalData

**License** Artistic-2.0

**biocViews** Software, Infrastructure, DataRepresentation

**Encoding** UTF-8

**RoxygenNote** 7.3.2

**VignetteBuilder** knitr

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**BugReports** <https://github.com/shbrief/OmicsMLRepoR/issues>

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availableTerms	<i>Extract all the terms used in a quired attribute/column</i>
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### Description

Extract all the terms used in a quired attribute/column

### Usage

```
availableTerms(attribute, db = "cMD")
```

### Arguments

attribute	A character (1). Name of the attribute/column you want to extract the terms used under.
db	A character(1). Currently, 'cMD' (curatedMetagenomicData) is only supported.

### Value

A data frame with two columns - existing values under the quired attribute ('allowedvalues' column) and their ontology term id ('ontology' column).

### Examples

```
availableTerms("age_group")
availableTerms("disease")
```

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getLongMetaTb	<i>Expands metadata with multiple values belong to a same attribute</i>
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### Description

Different from the 'getWideMetaTb' function, this function accepts multiple target columns ('targetCols'). The difference comes from that the target columns of this function are related, i.e., have the same number of elements separated by the 'delim', and the multiple values for each column belongs to the same column/attribute/field, i.e., no additional column name is required/provided.

### Usage

```
getLongMetaTb(meta, targetCols = NULL, delim = NULL)
```

### Arguments

meta	A data frame. Metadata table containing all treatment-related columns
targetCols	Optional. A character vector of column names to expand if present. Default is the name of all cBioPortal treatment-related columns.
delim	Optional. A character (1) of a delimiter used to separate multiple values in the metadata table.

### Value

A data frame of metadata expanded so that each individual treatment has its own row.

### Examples

```
data(mini_cmd)
lmeta <- getLongMetaTb(mini_cmd, "hla")
dim(mini_cmd)
dim(lmeta)

short_tb <- data.frame(
  ind = c("A", "B", "C", "D", "E"),
  aval = c("cat;dog", "chicken", "horse", "frog;pig", "snake"),
  cval = c(1, NA, 3, 4, 5),
  bval = c("red;blue", "yellow", "NA", "green;NA", "brown"))

getLongMetaTb(short_tb, c("aval", "bval"), delim = ";")
```

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getMetadata	<i>Download a harmonized metadata table</i>
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### Description

Download a harmonized metadata table

**Usage**

```
getMetadata(database = NULL, load = TRUE)
```

**Arguments**

database	Name of the database to get the metadata from. Currently, there are two available options. <ul style="list-style-type: none"> <li>• cmd : metadata for curatedMetagenomicData</li> <li>• cBioPortal : metadata for cBioPortalData</li> </ul>
load	Default is TRUE. If it's set to FALSE, the metadata table is downloaded to cache but not loaded into memory.

**Value**

Curated metadata table or file cache location, if 'load = FALSE'.

**Examples**

```
cmd <- getMetadata("cmd")
```

---

```
getNarrowMetaTb
```

*Collapse values from multiple columns into one*

---

**Description**

Collapse values from multiple columns into one

**Usage**

```
getNarrowMetaTb(
  meta,
  newCol = NULL,
  targetCols = NULL,
  sep = ":",
  delim = ";",
  remove = TRUE,
  na.rm = TRUE,
  sort = TRUE
)
```

**Arguments**

meta	A data frame.
newCol	A character (1). Name of the new column to store collapsed values.
targetCols	A character vector. Names of the columns to be collapsed into one column.
sep	A character (1). Delimiter used to concatenate column name and its value. Default is double colons, '::'.
delim	A character(1). Separator to use between values/columns. Default is ';'.

remove	With the default, 'TRUE', this function will remove input columns from output data frame.
na.rm	With the default, 'TRUE', missing values will be removed prior to uniting each value.
sort	With the default, 'TRUE', the united columns will be ordered alphabetically.

### Value

A data frame where target columns (targetCols) are collapsed into a single column. The original column name and its value are concatenated with the 'sep' input and the column:value pairs are separated by the 'delim' input. Target columns will be merged in the alphabetical order of their names.

### Examples

```
wide_tb <- data.frame(fruit = c("apple", "banana", "pear", "watermelon",
                              "grape"),
                    shape = c("round", "long", NA, "round", NA),
                    color = c("red", "yellow", NA, "green", "purple"),
                    size = c("medium", "medium", NA, "large", "small"))
getNarrowMetaTb(wide_tb,
                newCol = "feature",
                targetCols = c("color", "shape", "size"),
                sep = ":", delim = ";")
```

---

getOntoInfo

*Query Ontology Lookup Service (OLS)*


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### Description

Extract identical or similar ontology terms across different ontologies

### Usage

```
getOntoInfo(query, ontology = "", exact = FALSE, rows = 20)
```

### Arguments

query	A character (1) containing the search query, either a term label or term id.
ontology	A character vector defining the ontology to be queried. Default is the empty character, to search all ontologies.
exact	A logical (1) defining if OLS search is restricted to exact matches. Defaults is 'FALSE'.
rows	An integer (1) defining the number of query returns. Default is 20L. Maximum number of values returned by the server is 1000.

### Value

A tibble containing ontology term label and description

**Examples**

```

getOntoInfo("NCIT:C4872")
getOntoInfo("NCIT:C4872", ontology = c("EFO", "MONDO"))
getOntoInfo("Skin Infection")
getOntoInfo("Sitagliptin", ontology = c("Chebi", "apple"))

## Multiple query values
getOntoInfo("plasma,membrane")
getOntoInfo(c("plasma", "membrane"))

```

---

<code>getShortMetaTb</code>	<i>Compresses expanded metadata columns to one row per sample</i>
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---

**Description**

Compresses expanded metadata columns to one row per sample

**Usage**

```
getShortMetaTb(meta, idCols = NULL, targetCols = NULL, delim = "<>")
```

**Arguments**

<code>meta</code>	A data frame with expanded treatment columns.
<code>idCols</code>	Optional. A character vector of columns that identify single samples, such as 'curation_id' and 'sampleId'. Defaults to standard ID columns.
<code>targetCols</code>	Optional. A character vector of columns to compress if present. Default is names of all cBioPortal treatment-related columns.
<code>delim</code>	Optional. A delimiter string. Default is '<>'.

**Value**

A data frame where each sample gets a single row

**Examples**

```

data(mini_cmd)
lmeta <- getLongMetaTb(mini_cmd, "hla")
res <- getShortMetaTb(lmeta, targetCols = "hla")
dim(res) # 200 x 3 table

long_tb <- data.frame(ind = c("A", "A", "B", "C", "D", "D", "E"),
                     aval = c("cat", "dog", "chicken", "horse",
                              "frog", "pig", "snake"),
                     cval = c(1, 1, NA, 3, 4, 4, 5),
                     bval = c("red", "blue", "yellow", NA, "green",
                              NA, "brown"))
getShortMetaTb(long_tb, idCols = "ind", targetCols = c("aval", "bval"))

```

---

getWideMetaTb	<i>Create individual columns for different attributes stored in one column</i>
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---

### Description

The values stored in one column should include their potential column names to use this function.

### Usage

```
getWideMetaTb(meta, targetCol = NULL, sep = ":", delim = "<>", remove = TRUE)
```

### Arguments

meta	A data frame.
targetCol	A character. Names of the column whose contents are exposed as individual columns. Multiple attributes should be separated by the 'sep' and the column name and its value should be separated by the provided 'delim'.
sep	A character (1). Delimiter used to concatenate column name and its value. Default is double colons, '::'.
delim	A character(1). Separator used between values. Default '<>'.
remove	If 'TRUE', remove input columns from output data frame.

### Value

A data frame where the contents under 'targetCol' is split into individual columns in an alphabetical order. Data type of the expanded columns is all character.

### Examples

```
## Narrow-table example
narrow_tb <- data.frame(fruit = c("apple", "banana", "pear", "watermelon",
                                "grape"),
                      feature = c("color:red;shape:round;size:medium",
                                   "color:yellow;shape:long;size:medium",
                                   "color:brown;shape:NA;size:NA",
                                   "color:green;shape:round;size:large",
                                   "color:purple;shape:NA;size:small"))
getWideMetaTb(narrow_tb, targetCol = "feature", sep = ":", delim = "<>")

## Narrow-table example with missing columns
narrow_tb2 <- data.frame(fruit = c("apple", "banana", "pear",
                                   "watermelon", "grape"),
                      feature = c("color:red;shape:round;size:medium",
                                   "color:yellow;shape:long;size:medium",
                                   NA,
                                   "color:green;size:large",
                                   "color:purple;shape:NA;size:small"))
getWideMetaTb(narrow_tb2, targetCol = "feature", sep = ":", delim = "<>")
```

---

get_ontologies	<i>Extract ontology from the ontology term ids</i>
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---

### Description

Extract ontology from the ontology term ids

### Usage

```
get_ontologies(terms, delim = ":")
```

### Arguments

terms	A character vector
delim	A character. Delimiter between ontology and its id. Default is ':'.

### Value

A character vector containing the ontology names of the input 'terms'. The length of this is same as the 'terms' input.

### Examples

```
terms <- c("HP:0001824", "MONDO:0010200", "NCIT:C122328", "4471000175100")
get_ontologies(terms = terms)
```

---

merge_vectors	<i>Custom function to merge vectors</i>
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### Description

This function is designed for a group of, collapsible metadata attributes (e.g., 'biomarker' for curatedMetagenomicData).

### Usage

```
merge_vectors(base, update, sep = ":", delim = ";")
```

### Arguments

base	A character. A space-holder version of the key:value concatenates (e.g., 'column1:NA;column2:NA;column3:NA')
update	A character. The target string to be compared and filled with 'base' if there is missing pairs. (e.g., 'column1:value1;column3:value3')
sep	A character string to separate the column name and value. Default is ':'.
delim	A character string to separate the column:value pairs. Default is ';'.



**Value**

A character updated the target string ('update') to follow the reference string ('base').

**Examples**

```
x <- "color:NA;shape:NA;size:NA"
y <- "color:green;size:large"
merge_vectors(x, y)
```

---

mini\_cmd

*A subset of cMD metadata*

---

**Description**

A subset of curated version of 'sampleMetadata' from the curatedMetagenomicData (cMD, ver.3.8.0) package.

**Usage**

```
mini_cmd
```

**Format**

A data frame with 200 samples and 3 columns ('curation\_id', 'hla', and 'package')

**Author(s)**

Sehyun Oh <shbrief@gmail.com>

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OmicsMLRepoR

*OmicsMLRepoR: A package for browsing harmonized metadata through ontology*

---

**Description**

The OmicsMLRepoR package provides functions to browse the harmonized metadata created under the OmicsMLRepo project. It supports data navigation if the metadata incorporates ontology.

**Key Functions**

- [getMetadata](#): Download a harmonized metadata table
- [tree\\_filter](#): Find samples including the queried terms and their descendants

**Additional Information**

For more detailed information, see the vignette: `vignette("Quickstart", package = "OmicsMLRepoR")`

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ontoTreePlot	<i>Plot ontology tree</i>
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**Description**

Plot ontology tree

**Usage**

```
ontoTreePlot(term, display = c("Term", "Text"))
```

**Arguments**

term	A character (1). Ontology term id (obo_id)
display	A character (1) specifying a node labeling option. Two available options are 'Term' for ontology term or IRI (Internationalized Resource Identifier) and 'Text' for the label or preferred name.

**Value**

A ontology tree plot. All the terms used in the output plot are ancestors of the queried term, so the queried term is the tip.

**Examples**

```
ontoTreePlot("NCIT:C2852", "Term")
```

---

sample_metadata	<i>sample_metadata</i>
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---

**Description**

A small data table to demonstrating the data reshaping functions in OmicsMLRepoR.

**Usage**

```
sample_metadata
```

**Format**

A data frame with 4 rows and 7 columns

**Author(s)**

Sehyun Oh <shbrief@gmail.com>

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tree_filter	<i>Keep rows that include the queried terms and their descendants</i>
-------------	---

---

### Description

Similar to [filter](#) function, while its filtering includes descendants and synonyms of the query term in addition to ontology terms and ids identical or similar to the query term across different ontologies collected through OLS search.

### Usage

```
tree_filter(.data, col, query, logic = "OR", delim = NULL)
```

### Arguments

.data	A data frame
col	A character (1). Column name to filter by.
query	A character vector containing words or ids to be used in the ontology search
logic	A character (1). Operator used to determine filtering method. Values allowed: "AND", "OR", "NOT". Defaults to "OR"
delim	A character (1) used to separate multiple values. If your '.data' input is obtained from getMetadata function, this input is automatically configured.

### Value

Data frame filtered by provided queries along with child terms in the specified column

### Examples

```
meta <- getMetadata("CMD")
tree_filter(meta, disease, c("pancreatic disease", "cancer"))
```

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