

Package ‘pubchem.bio’

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Title Biologically Informed Metabolomic Libraries from 'PubChem'

Version 1.0.5

Description All 'PubChem' compounds are downloaded to a local computer, but for each compound, only partial records are used. The data are organized into small files referenced by 'PubChem' CID. This package also contains functions to parse the biologically relevant compounds from all 'PubChem' compounds, using biological database sources, pathway presence, and taxonomic relationships. Taxonomy is used to generate a lowest common ancestor taxonomy ID (NCBI) for each biological metabolite, which then enables creation of taxonomically specific metabolome databases for any taxon.

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Encoding UTF-8

Imports foreach, doParallel, R.utils, data.table, dplyr, redk, stringr, curl, MetaboCoreUtils, CHNOSZ, CompoundDb, RSQLite, magrittr

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 build.cid.lca

build.cid.lca

Description

utilizes downloaded and properly formatted local pubchem data created by 'get.pubchem.ftp' as input to generate a relationship between pubchem CID and the lowest common ancestor NCBI taxid

Usage

```

build.cid.lca(
  pc.directory = NULL,
  tax.sources = "LOTUS - the natural products occurrence database",
  use.pathways = TRUE,
  use.conservd.pathways = FALSE,
  threads = 8,
  cid.taxid.object = NULL,
  taxid.hierarchy.object = NULL,
  cid.pwid.object = NULL,
  min.taxid.table.length = 3,
  output.directory = NULL
)

```

Arguments

`pc.directory` directory from which to load pubchem .Rdata files. alternatively provide `cid.taxid.object`, `taxid.hierarchy.object`, and `cid.pwid.object` as `data.table` R objects.

`tax.sources` vector. which taxonomy sources should be used? defaults to `c("LOTUS - the natural products occurrence database", "The Natural Products Atlas", "KNAP-SAcK Species-Metabolite Database", "Natural Product Activity and Species Source (NPASS)")`.

`use.pathways` logical. default = TRUE, should pathway data be used in building lowest common ancestor, when taxonomy is associated with a pathway?

`use.conservd.pathways` logical. default = FALSE, should 'conserved' pathways be used? when false, only pathways with an assigned taxonomy are used.

`threads` integer. number of threads to use when finding lowest common ancestor. parallel processing via `DoParallel` and `foreach` packages.

`cid.taxid.object`
R `data.table`, generally produced by `get.pubchem.ftp`; preferably, define `pc.directory`

`taxid.hierarchy.object`
R `data.table`, generally produced by `get.pubchem.ftp`; preferably, define `pc.directory`

`cid.pwid.object`
R `data.table`, generally produced by `get.pubchem.ftp`; preferably, define `pc.directory`

`min.taxid.table.length`
integer. when there are few taxa reported to synthesize a particular compound, and those few taxa are spread widely across biology, the LCA concept breaks down. This value controls the decision as to whether to determine LCA within taxonomic ranks, rather within the full taxonomy hierarchy. see details.

`output.directory`
directory to which the `pubchem.bio` database is saved. If NULL, will try to save in `pc.directory` (if provided). If both directories are NULL, not saved, only returned as in memory

Details

utilizes downloaded and properly formatted local pubchem data created by 'get.pubchem.ftp' function

Some metabolism is highly conserved - all species perform those reactions. Other metabolism is highly specific - there is one known species to produce that metabolite. Sometimes, it is in between. The lowest common ancestor approach allows us to analyze these patterns and put them to use to generalize metabolites for metabolomics across species.

Biology is more complex than that though. Natural products are often reported as being synthesized by an organism which is in symbiosis with a second organism. The taxonomic assignment is sometimes both organisms, even if neither would create that product in isolation, or if only one is actually capable of producing that metabolite. In these situations, the LCA approach can break down. For example, if a bacteria is in symbiosis with an algae, and each is listed as producing the metabolite, the LCA will be assigned as '1' - the root of all biology, since we have to go back to the base of the taxonomic tree to find the common taxonomic ancestor of prokaryotes and eukaryotes. In this example, there are two unique species, genera, families, orders, etc listed in the full taxonomic hierarchy for this metabolite.

The 'min.unique.taxid.ct' variable controls sensitivity to this phenomenon in assigning LCA. The number of unique taxa which are mapped to each metabolite varies by taxonomic level. It may map to two species, but only one genus. In that case, the genus is assigned as the LCA. However, if the metabolite maps to two unique species, two unique genera, two unique families, two unique kingdoms, and one unique domain, we should ask ourselves whether this sparse pattern supports that this metabolite should be marked as 'conserved' or 'primary.' What makes more intuitive sense is to conclude that there may be extenuating circumstances which have resulted from unique biology. For example, Ceratodictyol B is reported from *Haliclona cymaeformis* and *Ceratodictyon spongiosum*, one of which is a red algal symbiont of the other. At each taxonomic level, there are either 0, 1, or 2 unique taxonomy IDs. 0 unique levels is uninteresting - that just reflects that there is no taxonomy assigned for those lineages at that level.

What is more interesting is the number of unique levels of the number of unique taxonomy IDs. In the case of Ceratodictyol B, the only other value is '2'. There are 2 unique taxonomy IDs at each level species, genus, order, class, and phylum. So there are five taxonomic levels that have exactly 2 unique taxonomy IDs, and there are no taxonomic levels which have more than 2 unique taxids. We will call this the taxid.ct.length, where the taxid.ct.table is the table of frequencies of the number of unique taxids at each taxonomic level. The length is the number of unique values when ignoring '0' or '1'. When the taxid.ct.length is less than or equal to min.taxid.table.length, the lca is calculated within the lowest taxonomic level that has the most frequent unique taxonomy ID count.

For the Ceratodictyol B example, this would mean that we would find that '2' was the most common number of unique taxids reported, so we find that the lowest taxonomic level which reports two unique taxids is 'species'. LCA is for assigned to those two species. If however, there were two *Ceratodictyon* spp reported, then the species level would have 3 unique taxids, and there would be 4 levels (rather than five) which have 2 unique taxids. The lowest taxonomic level with 2 unique taxids, the most frequent count observed, would now be 'genus', so LCA would be assigned for within each level of 'genus'. This would mean that the first LCA would be assigned to the *Ceratodictyon* genus, since there are multiple *Ceratodictyon* species reported, and then a second LCA would be assigned to the *Haliclona cymaeformis* species. Sorry it is so complicated. Life is complicated.

Value

nothing. will save to pc.directory as .Rdata file.

Author(s)

Corey Broeckling

Examples

```
data('cid.taxid', package = "pubchem.bio")
data('taxid.hierarchy', package = "pubchem.bio")
data('cid.pwid', package = "pubchem.bio")
cid.lca.out <- build.cid.lca(
  tax.sources = "LOTUS - the natural products occurrence database",
  use.pathways = FALSE,
  threads = 1, cid.taxid.object = cid.taxid,
  taxid.hierarchy.object = taxid.hierarchy,
  cid.pwid.object = cid.pwid)
head(cid.lca.out)
```

build.element.count *build.element.count*

Description

takes as input an pubchem.bio data.table (generally produced by 'build.pubchem.bio' or 'build.taxon.metabolome') and removes inorganic compounds (those without any carbon).

Usage

```
build.element.count(
  pc.directory = NULL,
  pubchem.bio.object = NULL,
  remove.homonuclear.molecules = TRUE,
  remove.inorganics = TRUE
)
```

Arguments

pc.directory directory from which to load pubchem .Rdata files

pubchem.bio.object
R data.table, generally produced by build.pubchem.bio; preferably, define pc.directory

remove.homonuclear.molecules
logical. default = TRUE. should molecules with one element (or fewer) be removed?

remove.inorganics
logical. default = TRUE. Should inorganic molecules (no carbon) be removed?

Details

utilizes downloaded and properly formatted local pubchem data created by 'build.pubchem.bio' function, removes inorganic compounds. note that for any formula which have charge noted '+' or '-', the charge is removed for tabulation of the element count.

Value

a data.table, same as input, except inorganics will have been removed.

Author(s)

Corey Broeckling

build.primary.metabolome

build.primary.metabolome

Description

utilizes downloaded and properly formatted local pubchem data created by 'get.pubchem.ftp' function to filter a dataset created by 'build.pubchem.bio' function

Usage

```
build.primary.metabolome(
  pc.directory = NULL,
  get.properties = FALSE,
  threads = 8,
  db.name = "primary.metabolome",
  rcdk.desc = c("org.openscience.cdk.qsar.descriptors.molecular.XLogPDescriptor",
    "org.openscience.cdk.qsar.descriptors.molecular.AcidicGroupCountDescriptor",
    "org.openscience.cdk.qsar.descriptors.molecular.BasicGroupCountDescriptor",
    "org.openscience.cdk.qsar.descriptors.molecular.TPSADescriptor"),
  pubchem.bio.object = NULL,
  output.directory = NULL,
  keep.primary.only = TRUE,
  min.tax.ct = 3
)
```

Arguments

pc.directory	directory from which to load pubchem .Rdata files
get.properties	logical. if TRUE, will return rcdk calculated properties: XLogP, TPSA, HBond-DonorCount and HBondAcceptorCount.
threads	integer. how many threads to use when calculating rcdk properties. parallel processing via DoParallel and foreach packages.

Usage

```

build.pubchem.bio(
  pc.directory = NULL,
  use.bio.sources = TRUE,
  bio.sources = c("Metabolomics Workbench", "Human Metabolome Database (HMDB)", "ChEBI",
    "LIPID MAPS", "MassBank of North America (MoNA)"),
  use.pathways = TRUE,
  pathway.sources = NULL,
  use.taxid = TRUE,
  taxonomy.sources = NULL,
  use.parent.cid = TRUE,
  use.parent.when.charged = FALSE,
  remove.salts = TRUE,
  remove.inorganics = FALSE,
  mw.range = c(50, 2000),
  get.properties = TRUE,
  threads = 8,
  rcdk.desc = c("org.openscience.cdk.qsar.descriptors.molecular.XLogPDescriptor",
    "org.openscience.cdk.qsar.descriptors.molecular.AcidicGroupCountDescriptor",
    "org.openscience.cdk.qsar.descriptors.molecular.BasicGroupCountDescriptor",
    "org.openscience.cdk.qsar.descriptors.molecular.TPSADescriptor"),
  cid.lca.object = NULL,
  cid.sid.object = NULL,
  cid.pwid.object = NULL,
  cid.parent.object = NULL,
  cid.taxid.object = NULL,
  cid.formula.object = NULL,
  cid.smiles.object = NULL,
  cid.inchikey.object = NULL,
  cid.inchi.object = NULL,
  cid.monoisotopic.mass.object = NULL,
  cid.title.object = NULL,
  cid.cas.object = NULL,
  cid.pmid.ct.object = NULL,
  output.directory = NULL
)

```

Arguments

- pc.directory** directory from which to load pubchem .Rdata files. alternatively, provide R data.tables for ALL *cid.property.object* options defined below.
- use.bio.sources** logical. If TRUE (default) use the bio.source vector of sources, incorporating all CIDs from those bio databases.
- bio.sources** vector of source names from which to extract pubchem CIDs. all can be found here: <https://pubchem.ncbi.nlm.nih.gov/sources/>, but can additionally use "PubChemLite" as a datasource. defaults to c("Metabolomics Workbench", "Human Metabolome Database (HMDB)", "ChEBI", "LIPID MAPS", "MassBank

cid.smiles.object
R data.table, generally produced by get.pubchem.ftp; preferably, define pc.directory

cid.inchikey.object
R data.table, generally produced by get.pubchem.ftp; preferably, define pc.directory

cid.inchi.object
R data.table, generally produced by get.pubchem.ftp; preferably, define pc.directory

cid.monoisotopic.mass.object
R data.table, generally produced by get.pubchem.ftp; preferably, define pc.directory

cid.title.object
R data.table, generally produced by get.pubchem.ftp; preferably, define pc.directory

cid.cas.object R data.table, generally produced by get.pubchem.ftp; preferably, define pc.directory

cid.pmid.ct.object
R data.table, generally produced by get.pubchem.ftp; preferably, define pc.directory

output.directory
directory to which the pubchem.bio database is saved. If NULL, will try to save in pc.directory (if provided), else not saved.

Details

utilizes downloaded and properly formatted local pubchem data created by 'get.pubchem.ftp' function

Value

a data frame containing pubchem CID, title, formula, monoisotopic molecular weight, inchikey, smiles, cas, optionally rcdk properties

Author(s)

Corey Broeckling

Examples

```
data('cid.sid', package = "pubchem.bio")
data('cid.pwid', package = "pubchem.bio")
data('cid.parent', package = "pubchem.bio")
data('cid.taxid', package = "pubchem.bio")
data('cid.formula', package = "pubchem.bio")
data('cid.smiles', package = "pubchem.bio")
data('cid.inchikey', package = "pubchem.bio")
data('cid.inchi', package = "pubchem.bio")
data('cid.monoisotopic.mass', package = "pubchem.bio")
data('cid.title', package = "pubchem.bio")
data('cid.cas', package = "pubchem.bio")
data('cid.pmid.ct', package = "pubchem.bio")
data('cid.lca', package = "pubchem.bio")
pc.bio.out <- build.pubchem.bio(use.pathways = FALSE, use.parent.cid = FALSE,
get.properties = FALSE, threads = 1,
cid.sid.object = cid.sid, cid.pwid.object = cid.pwid,
```

```

cid.parent.object = cid.parent, cid.taxid.object = cid.taxid,
cid.formula.object = cid.formula, cid.smiles.object = cid.smiles,
cid.inchikey.object = cid.inchikey, cid.inchi.object = cid.inchi,
cid.monoisotopic.mass.object = cid.monoisotopic.mass,
cid.title.object = cid.title, cid.cas.object = cid.cas,
cid.pmid.ct.object = cid.pmid.ct, cid.lca.object = cid.lca)
head(pc.bio.out)

```

```
build.taxon.metabolome
```

```
build.taxon.metabolome
```

Description

utilizes downloaded and properly formatted local pubchem data created by 'get.pubchem.ftp' function to filter a dataset created by 'build.pubchem.bio' function

Usage

```

build.taxon.metabolome(
  pc.directory = NULL,
  taxid = c(),
  get.properties = FALSE,
  full.scored = TRUE,
  keep.scored.only = FALSE,
  aggregation.function = max,
  threads = 8,
  db.name = "custom.metabolome",
  rcdk.desc = c("org.openscience.cdk.qsar.descriptors.molecular.XLogPDescriptor",
    "org.openscience.cdk.qsar.descriptors.molecular.AcidicGroupCountDescriptor",
    "org.openscience.cdk.qsar.descriptors.molecular.BasicGroupCountDescriptor",
    "org.openscience.cdk.qsar.descriptors.molecular.TPSADescriptor"),
  pubchem.bio.object = NULL,
  cid.lca.object = NULL,
  taxid.hierarchy.object = NULL,
  output.directory = NULL
)

```

Arguments

pc.directory	directory from which to load pubchem .Rdata files
taxid	integer vector of integer NCBI taxonomy IDs. i.e. c(9606, 1425170) for Homo sapiens and Homo heidelbergensis.
get.properties	logical. if TRUE, will return rcdk calculated properties: XLogP, TPSA, HBond-DonorCount and HBondAcceptorCount.

Examples

```
data('cid.lca', package = "pubchem.bio")
data('pubchem.bio', package = "pubchem.bio")
data('taxid.hierarchy', package = "pubchem.bio")
my.taxon.db <- build.taxon.metabolome(
  pubchem.bio.object = pubchem.bio,
  cid.lca.object = cid.lca, taxid.hierarchy.object = taxid.hierarchy,
  get.properties = FALSE, threads = 1, taxid = c(1))
head(my.taxon.db)
```

cid.accurate.mass	<i>cid.accurate.mass.rda</i>
-------------------	------------------------------

Description

A subset of the full cid.accurate.mass, for example code

Format

data.table, stored in .rda format

Source

subset of cid.accurate.mass file from get.pubchem.ftp

cid.cas	<i>cid.cas.rda</i>
---------	--------------------

Description

A subset of the full cid.cas, for example code

Format

data.table, stored in .rda format

Source

subset of cid.cas file from get.pubchem.ftp

cid.formula	<i>cid.formula.rda</i>
-------------	------------------------

Description

A subset of the full cid.formula, for example code

Format

data.table, stored in .rda format

Source

subset of cid.formula file from get.pubchem.ftp

cid.inchi	<i>cid.inchi.rda</i>
-----------	----------------------

Description

A subset of the full cid.inchi, for example code

A subset of the full cid.inchi, for example code

Format

data.table, stored in .rda format

data.table, stored in .rda format

Source

subset of cid.inchi file from get.pubchem.ftp

subset of cid.inchi file from get.pubchem.ftp

cid.inchikey	<i>cid.inchikey.rda</i>
--------------	-------------------------

Description

A subset of the full cid.inchikey, for example code

Format

data.table, stored in .rda format

Source

subset of cid.inchikey file from get.pubchem.ftp

cid.lca	<i>cid.lca.rda</i>
---------	--------------------

Description

A subset of the full cid.lca, for example code

Format

data.table, stored in .rda format

Source

subset of cid.lca file from get.pubchem.ftp

cid.mesh.function	<i>cid.mesh.function.rda</i>
-------------------	------------------------------

Description

A subset of the full cid.mesh, for example code

Format

data.table, stored in .rda format

Source

subset of cid.mesh file from get.pubchem.ftp

cid.mesh.name	<i>cid.mesh.name.rda</i>
---------------	--------------------------

Description

A subset of the full cid.mesh.name, for example code

Format

data.table, stored in .rda format

Source

subset of cid.mesh.name file from get.pubchem.ftp

cid.monoisotopic	<i>cid.monoisotopic.mass.rda</i>
------------------	----------------------------------

Description

A subset of the full cid.monoisotopic, for example code

Format

data.table, stored in .rda format

Source

subset of cid.monoisotopic file from get.pubchem.ftp

cid.monoisotopic.mass	<i>cid.monoisotopic.mass.rda</i>
-----------------------	----------------------------------

Description

A subset of the full cid.monoisotopic.mass, for example code

Format

data.table, stored in .rda format

Source

subset of cid.accurate.mass file from get.pubchem.ftp

cid.parent	<i>cid.parent.rda</i>
------------	-----------------------

Description

A subset of the full cid.parent, for example code

Format

data.table, stored in .rda format

Source

subset of cid.parent file from get.pubchem.ftp

cid.pmid	<i>cid.pmid.rda</i>
----------	---------------------

Description

A subset of the full cid.pmid, for example code

Format

data.table, stored in .rda format

Source

subset of cid.pmid file from get.pubchem.ftp

cid.pmid.ct	<i>cid.pmid.ct.rda</i>
-------------	------------------------

Description

A subset of the full cid.pmid.ct, for example code

Format

data.table, stored in .rda format

Source

subset of cid.pmid.ct file from get.pubchem.ftp

cid.preferred	<i>cid.preferred.rda</i>
---------------	--------------------------

Description

A subset of the full cid.preferred, for example code

Format

data.table, stored in .rda format

Source

subset of cid.preferred file from get.pubchem.ftp

cid.pwid	<i>cid.pwid.rda</i>
----------	---------------------

Description

A subset of the full cid.pwid, for example code

Format

data.table, stored in .rda format

Source

subset of cid.pwid file from get.pubchem.ftp

cid.sid	<i>cid.sid.rda</i>
---------	--------------------

Description

A subset of the full cid.sid, for example code

Format

data.table, stored in .rda format

Source

subset of cid.sid file from get.pubchem.ftp

cid.smiles	<i>cid.smiles.rda</i>
------------	-----------------------

Description

A subset of the full cid.smiles, for example code

Format

data.table, stored in .rda format

Source

subset of cid.smiles file from get.pubchem.ftp

cid.synonym	<i>cid.synonym.rda</i>
-------------	------------------------

Description

A subset of the full cid.synonym, for example code

Format

data.table, stored in .rda format

Source

subset of cid.synonym file from get.pubchem.ftp

cid.taxid	<i>cid.taxid.rda</i>
-----------	----------------------

Description

A subset of the full cid.taxid, for example code

Format

data.table, stored in .rda format

Source

subset of cid.taxid file from get.pubchem.ftp

cid.title	<i>cid.title.rda</i>
-----------	----------------------

Description

A subset of the full cid.title, for example code

Format

data.table, stored in .rda format

Source

subset of cid.title file from get.pubchem.ftp

export.CompoundDb *export.CompoundDb*

Description

export pubchem.bio pc.bio syle data.table to a CompoundDb database

Usage

```
export.CompoundDb(
  pc.bio.object = NULL,
  pc.directory = NULL,
  path = NULL,
  dbFile = NULL,
  source = NULL,
  source_version = NULL,
  url = NULL,
  source_date = NULL,
  organism = NA_character_,
  get.synonyms = FALSE
)
```

Arguments

pc.bio.object	input data.table, generated from 'build.pubchem.bio' or 'build.taxon.metabolome' functions
pc.directory	directory from which to load pubchem .Rdata files. alternatively, provide R data.tables for ALL cid.property.object options defined below.
path	valid file path specifying the directory to which the database will be saved.
dbFile	valid file name - use .sqlite extension. if NULL, will set to 'pubchem.bio.sqlite'
source	default = NULL. if NULL, will set to 'pubchem.bio'
source_version	default = NULL. if NULL, will set to pubchem.bio version.
url	default = NULL. if NULL, will set to "https://cran.r-project.org/web/packages/pubchem.bio/index.html"
source_date	default = NULL. if NULL, will set to current date via format(Sys.Date(), format = "%Y-%m-%d")
organism	character. default = NA_character_. if this is a species specific database, set to species name or NCBI taxonomy ID number.
get.synonyms	logical. default = FALSE. When TRUE, will load the 'cid.synonyms.Rdata' file from the pc.directory, which must be defined. database product will have all synonyms included. If FALSE, synonyms will be an empty list.

Details

takes output from 'build.pubchem.bio' or 'build.taxon.metabolome' functions, reformatting, and converting to a CompoundDb object and database. see vignette('create-compounddb', package = 'CompoundDb') for more details on the CompoundDb package

Value

'CompDb' from CompoundDb package. see vignette('create-compounddb', package = 'CompoundDb')

Author(s)

Corey Broeckling

export.msfinder	<i>export.msfinder</i>
-----------------	------------------------

Description

export pubchem.bio pc.bio syle data.table to format suitable for MSFinder input.

Usage

```
export.msfinder(pc.bio.object = NULL, export.file.name = NULL)
```

Arguments

pc.bio.object input data.table, generated from 'build.pubchem.bio' or 'build.taxon.metabolome' functions
export.file.name valid file path and name. Extension should be listed as '.tsv'.

Details

takes output from 'build.pubchem.bio' or 'build.taxon.metabolome' functions, reformatting, and exporting to input format suitable for MSFinder.

Value

nothing - file written to disk.

Author(s)

Corey Broeckling

Arguments

pc.bio.object input data.table, generated from 'build.pubchem.bio' or 'build.taxon.metabolome' functions
 export.file.name valid file path and name. Extension should be listed as '.tsv'.

Details

takes output from 'build.pubchem.bio' or 'build.taxon.metabolome' functions, reformatting, and exporting to input format suitable for Sirius.

Value

nothing - file written to disk.

Author(s)

Corey Broeckling

<code>get.pubchem.ftp</code>	<i>get.pubchem.ftp</i>
------------------------------	------------------------

Description

first step to building a local selective, biologically focused, pubchem data repository focused on metabolomics informatics

Usage

```
get.pubchem.ftp(  
  pc.directory = NULL,  
  timeout = 50000,  
  rm.tmp.files = TRUE,  
  threads = 2  
)
```

Arguments

pc.directory character. directory to which data will be saved
 timeout numeric. timeout setting for FTP download. setting options(timeout) value too small will generate errors for large files. default = 50000.
 rm.tmp.files logical. should temporary files be removed after completion of download and parsing? Default = TRUE.
 threads integer. the number of parallel threads to be used by foreach %dopar% during processing of taxonomy hierarchy data.

Details

this function downloads and unzips files from pubchem and NCBI taxonomy FTP sites as a first step in building a local metabolomics repository.

Value

nothing. all data are saved to disk for later loading

Author(s)

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Examples

```
## Not run:
my.dir <- "C:/Temp/20250725"
# or some other valid directory.
# this will be created assuming 'C:/Temp' exists.
get.pubchem.ftp(
  pc.directory = my.dir,
  timeout = 50000,
  rm.tmp.files = TRUE
)

## End(Not run)
```

pc.bio.subset

pc.bio.subset.rda

Description

A small dataset of a pubchem.bio metabolome scored by taxon, for inclusion in vignette

Format

data.table, stored in .rda format

Source

pubchem.bio data.table output derived from build.pubchem.bio function

pubchem.bio	<i>pubchem.bio.rda</i>
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Description

A subset of a full pubchem.bio biological metabolome, for example code

Format

data.table, stored in .rda format

Source

subset of pubchem.bio file from build.pubchem.bio function

sub.taxid.hierarchy	<i>sub.taxid.hierarchy.rda</i>
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Description

A small dataset of a the taxonomy hierarchy, for inclusion in vignette

Format

data.table, stored in .rda format

Source

pubchem.bio data.table output from NCBI Taxonomy data

taxid.hierarchy	<i>taxid.hierarchy.rda</i>
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Description

A subset of the full taxid.hierarchy, for example code

Format

data.table, stored in .rda format

Source

pubchem.bio data.table output from NCBI Taxonomy data

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