

Package ‘Onassis’

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Title OnASSIs Ontology Annotation and Semantic Similarity software

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Description

A package that allows the annotation of text with ontology terms (mainly from OBO ontologies) and the computation of semantic similarity measures based on the structure of the ontology between different annotated samples.

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Depends R (>= 3.5), rJava, OnassisJavaLibs

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annotate annotate

Description

This method annotates the entities contained in a data frame with the concepts from a specific dictionary.

Usage

```
annotate(input = NA, dictType = NA, dictionary = NA, ...)
```

```
## S4 method for signature 'data.frame,character,character'
annotate(input,
  dictType = "OBO", dictionary = NA, dictoutdir = getwd(),
  d_synonymtype = "EXACT", taxID = 0, annot_out = getwd(),
  paramValueIndex = NA, SearchStrategy = "CONTIGUOUS_MATCH",
  CaseMatch = "CASE_INSENSITIVE", Stemmer = "NONE",
  StopWords = "NONE", OrderIndependentLookup = "ON",
  FindAllMatches = "YES", e_synonymtype = "ALL",
  multipleDocs = FALSE, disease = FALSE)
```

Arguments

input	A data frame where the first column is the ID of the sample or document to annotate
dictType	the type of input dictionary OBO A dictionary that has been created by An OBO file ENTREZ Entrez genes dictionary TARGET Entrez genes dictionary, Histone marks and Histone modifications CMDICT A previously created dictionary file in the Conceptmapper XML format
dictionary	The local OBO/OWL ontology to be converted into an XML Conceptmapper dictionary or the URL to download the file. If NA is passed and the dictType parameter is not the default OBO then the method tries to download the corresponding dictionary from the available repositories. For ENTREZ and TARGET dictionary types a file named gene_info.gz can be automatically downloaded from ftp://ncbi.nlm.nih.gov/gene/data/gene_info.gz if its path is not provided by the user in this parameter. Alternatively an annotation package of the type Org.xx.eg.db from Bioconductor can be used. In this case the gene identifiers and their alternative names will be retrieved from the annotation database without the need of downloading a gene_info file.
...	Optional parameters
dictoutdir	Optional parameter to specify the location where the Conceptmapper dictionary file will be stored. Defaults to current working directory.

d_synonymtype	Optional parameter to specify the type of synonyms to consider when building the dictionary for Conceptmapper. For further detail http://owcollab.github.io/oboformat/doc/obo-syntax.html . Default: EXACT EXACT ALL
taxID	the taxonomy identifier of the organism when the dictType = 'ENTREZ' or 'TARGET' and the dictionary parameter refers to a gene_info.gz file. If 0 all the taxonomies will be included in the new dictionary.
annot_out	The path of the output directory where Conceptmapper annotation files will be stored
paramValueIndex	An integer value to index the 576 parameter combinations
SearchStrategy	The matching strategy for finding concepts in the input text <ul style="list-style-type: none"> • CONTIGUOUS_MATCH Longest match of contiguous tokens within enclosing span • SKIP_ANY_MATCH Longest match of not-necessarily contiguous tokens • SKIP_ANY_MATCH_ALLOW_OVERLAP Longest match of not-necessarily contiguous tokens, overlapping matches are allowed
CaseMatch	<ul style="list-style-type: none"> • CASE_IGNORE Fold everything to lowercase for matching • CASE_INSENSITIVE Fold only tokens with initial caps to lowercase • CASE_FOLD_DIGITS Fold all (and only) tokens with a digit • CASE_SENSITIVE Perform no case folding
Stemmer	<ul style="list-style-type: none"> • BIOLEMMATIZER A stemmer specific for biomedical literature • PORTER A stemmer that removes the commoner morphological and inflexional endings from words in English • NONE No word stemming
StopWords	<ul style="list-style-type: none"> • PUBMED A list of stop words obtained analyzing Pubmed papers • NONE No stop words
OrderIndependentLookup	<ul style="list-style-type: none"> • ON Ordering within span is ignored (i.e. 'Breast cancer' would equal 'Cancer breast') • OFF Ordering is taken into consideration
FindAllMatches	<ul style="list-style-type: none"> • YES All the matches within the span are found • NO Only the longest match within the span will be returned
e_synonymtype	The type of synonyms for the EntityFinder <ul style="list-style-type: none"> • EXACT_ONLY Only exact synonyms are considered • ALL All synonym types are included
multipleDocs	TRUE when multiple documents are loaded from a single file with each row representing a document. The file should have two columns. The first for the unique document identifier and the second for the textual descriptions
disease	A logical value set to TRUE if the annotation requires the 'Healthy' condition to be found.

Value

instance of class [Onassis-class](#) with annotated entities

Examples

```
geo_chip <- readRDS(system.file('extdata', 'vignette_data',
                              'GEO_human_chip.rds', package='Onassis'))

obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
onassis_results <- annotate(geo_chip, 'OBO', dictionary=obo)
entities <- entities(onassis_results)
entities <- entities[sample(nrow(entities), 30),]
```

annotateDF	annotateDF
------------	------------

Description

Method to find concepts from a Conceptmapper Dictionary of type [CMdictionary](#) contained in a given data frame, with a specified configuration of type [CMoptions](#). This is a method of the [EntityFinder-class](#)

Usage

```
annotateDF(object, descr_df, outDir = tempdir(), configOpt, cmDict)

## S4 method for signature 'EntityFinder,data.frame,character,CMoptions'
annotateDF(object,
           descr_df, outDir = tempdir(), configOpt, cmDict)
```

Arguments

object	Instance of class EntityFinder-class
descr_df	the table of text to annotate. The data frame should have identifiers in the first column and descriptions or text in the rest of the columns.
outDir	the output directory
configOpt	instance of class CMoptions-class
cmDict	Object of type CMdictionary-class containing the reference to a previously created Conceptmapper dictionary. Alternatively the path to a Conceptmapper xml file can be passed.

Value

A data frame of annotations containing the sample name, the id of the OBO concept, the corresponding name, the part of the text containing the annotation

Examples

```
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
dict <- CMdictionary(inputFileOrDb=obo, outputDir=getwd(), synonymType='ALL')
opts <- CMoptions()
ef <- new('EntityFinder')
methylation <- readRDS(system.file('extdata', 'vignette_data',
                                  'GEOmethylation.rds', package='Onassis'))
annotations <- annotateDF(ef, methylation[1:10, ], getwd(), opts, dict)
```

```
annotateTissueDisease  annotateTissueDisease
```

Description

annotateTissueDisease is a function to automatize the annotation process of tissues and diseases. It connects to the GEOmetadb through the `geo_metadb_path` parameter to retrieve the metadata of the samples provided in the `gsm_list` parameter. A dictionary for tissues/cell lines is built from the `tissue_obo` file provided as parameter. All the samples' metadata are annotated with tissue concepts from the `tissue_obo` and samples are clustered based on the semantic similarity of the defined semantic annotation sets. To reduce the number of tissue semantic sets, similar semantic sets are merged based on a given semantic similarity threshold provided in the `height_threshold` parameter. Within each semantic set, samples are annotated with disease concepts from the dictionary obtained from `disease_obo` parameter. For each disease the function retrieves the columns of the `score_matrix` to organize them in a list with tissues, diseases and scores.

Usage

```
annotateTissueDisease(geo_metadb_path, gsm_list, tissue_obo, disease_obo,
  outdir, height_threshold, score_matrix)
```

Arguments

<code>geo_metadb_path</code>	The full path of the directory where the GEOmetadb.sqlite file is stored
<code>gsm_list</code>	A list of GEO sample ids (GSM)s to annotate with tissue and disease concepts
<code>tissue_obo</code>	The obo ontology containing concepts to identify tissues/cell lines
<code>disease_obo</code>	The obo ontology containing concepts to identify diseases
<code>outdir</code>	The directory where the results will be stored
<code>height_threshold</code>	The percentage of clusters to merge based on the height of the dendrogram produced by the hclust method. Height_threshold is defined in the range [0, 1].
<code>score_matrix</code>	A matrix where rows represent units (GRanges or genes) and columns represent GSMs.

Value

A list of the tissue semantic sets defined by Onassis. For each tissue, a list of diseases and for each disease the columns of the `score_matrix` that were annotated with a given tissue and a given disease

Examples

```
if(!file.exists(file.path(getwd(), 'GEOmetadb.sqlite'))){
  message('To run this example please copy GEOmetadb.sqlite in your current working directory')
} else{
  geo_metadb_path <- getwd()
  score_matrix <- readRDS(system.file('extdata', 'score_matrix.rds', package='Onassis'))
  gsm_list <- colnames(score_matrix)
  tissue_obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
```

```

disease_obo <- system.file('extdata', 'sample.do.obo', package='OnassisJavaLibs')
outdir = getwd()
height_threshold <- 0.4
result_list <- annotateTissueDisease(geo_metadb_path, gsm_list, tissue_obo, disease_obo, outdir, height_
}

```

CaseMatch

CaseMatch

Description

Method to get and set the CaseMatch parameter of [COptions](#) instances

Usage

```

CaseMatch(x)

CaseMatch(x) <- value

## S4 method for signature 'COptions'
CaseMatch(x)

## S4 replacement method for signature 'COptions'
CaseMatch(x) <- value

```

Arguments

x	instance of class COptions
value	One of the following : <ul style="list-style-type: none"> • <code>CASE_IGNORE</code> Fold everything to lowercase for matching • <code>CASE_INSENSITIVE</code> Fold only tokens with initial caps to lowercase • <code>CASE_FOLD_DIGITS</code> Fold all (and only) tokens with a digit • <code>CASE_SENSITIVE</code> Perform no case folding

Value

The CaseMatch corresponding to the current options when used as get, the new COptions object with updated parameters when used to set.

Examples

```

opts <- COptions()
CaseMatch(opts)
opts <- COptions()
CaseMatch(opts) <- 'CASE_SENSITIVE'

```

 CMdictionary

 CMdictionary

Description

Constructor method for creating instances of class [CMdictionary-class](#). The created Conceptmapper dictionary will be stored as an XML file in the file system.

Usage

```
CMdictionary(inputFileOrDb = NULL, dictType = "OBO",
             outputDir = tempdir(), synonymType = "EXACT", taxID = 0,
             outputDirOp = TRUE)
```

Arguments

inputFileOrDb	The local OBO/OWL ontology to be converted into an XML Conceptmapper dictionary or the URL of a OBO/OWL file. If inputFileOrDb is NA and the dictType parameter is not the generic OBO then the method tries to download the corresponding dictionary from the available repositories. For ENTREZ and TARGET dictionary types a file named gene_info.gz will be automatically downloaded from ftp://ncbi.nlm.nih.gov/gene/data/gene_info.gz if a valid path is not provided by the user. Alternatively the name of an annotation package of the type Org.xx.eg.db from Bioconductor can be used. In this case the gene unique identifiers and their alternative identifiers will be retrieved from the annotation database without the need of downloading a gene_info file.
dictType	the type of input dictionary OBO A dictionary that has been created by An OBO file ENTREZ Entrez genes dictionary TARGET Entrez genes dictionary, Histone marks and Histone modifications CMDICT A previously created dictionary file in the Conceptmapper XML format
outputDir	the directory where the XML conceptmapper dictionary will be stored. Defaults to the tmp system's directory
synonymType	The type of synonyms to consider when building the dictionary for Conceptmapper. For further detail http://owcollab.github.io/oboformat/doc/obo-syntax.html . Default: EXACT EXACT ALL
taxID	the taxonomy identifier of the organism when the dictionary type is ENTREZ or TARGET. If 0 all the taxonomies will be included in the new dictionary.
outputDirOp	set to TRUE to clean the directory before creating the dictionary

Value

An object of type [CMdictionary-class](#) that can be used to annotate text with the [EntityFinder](#).

Examples

```
## Not run:
#' ##This might take some time to download the dictionary
dict <- CMdictionary(dictType = 'TARGET', inputFileOrDb='org.Hs.eg.db')

dict_file <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
dictionary <- CMdictionary(dictType='OBO', inputFileOrDb=dict_file)

## End(Not run)
```

CMdictionary-class *Class that stores a Conceptmapper dictionary*

Description

CMdictionary is a class that wraps a Conceptmapper ccp-nlp Java dictionary. Its methods allow the creation of a dictionary from OBO ontologies in OBO or OWL format. Different options to build the dictionary are available.

Details

The following methods can be applied to CMdictionary instances

```
dict_location
dict_location<-
dictInfo
dictInfo<-
dictRef
dictRef<-
```

To show the available dictionary types use the function `dictTypes`

Slots

`dict_location` The path of the created dictionary file

`dictRef` Reference to the java object representing the dictionary

`dictInfo` Information about how dictionary has been created. It is a list with the following fields

- Dictionary Type: The type of dictionary
 - OBOA dictionary that has been created by An OBO file
 - ENTREZEntrez genes dictionary
 - TARGETEntrez genes dictionary, Histone marks and Histone modifications
 - CMDICTA previously created dictionary file in the Conceptmapper XML format
- SynonymType: The type of synonyms to consider when building the dictionary for Conceptmapper. For further detail <http://owlcollab.github.io/oboformat/doc/obo-syntax.html>
 - EXACT
 - BROAD
 - NARROW

- RELATED
- ALL
- dictSource: The OBO/OWL dictionary file to convert to a Conceptmapper dictionary in case the type is OBO. The XML file in case the type is CMDICT
- taxID The NCBI taxon identifier for species to create the Entrez gene dictionary (e.g 9606 for *Mus musculus*)

Examples

```
dict <- new('CMdictionary')
```

CMoptions	CMoptions
-----------	-----------

Description

This constructor creates an object of type CMoptions

Usage

```
CMoptions(SearchStrategy = "CONTIGUOUS_MATCH",
           CaseMatch = "CASE_INSENSITIVE", Stemmer = "NONE",
           StopWords = "NONE", OrderIndependentLookup = "ON",
           FindAllMatches = "YES", SynonymType = "ALL", paramValueIndex = NA)
```

Arguments

- | | |
|------------------------|--|
| SearchStrategy | <p>The matching strategy for finding concepts in the input text</p> <ul style="list-style-type: none"> • CONTIGUOUS_MATCH Longest match of contiguous tokens within enclosing span • SKIP_ANY_MATCH Longest match of not-necessarily contiguous tokens • SKIP_ANY_MATCH_ALLOW_OVERLAP Longest match of not-necessarily contiguous tokens, overlapping matches are allowed |
| CaseMatch | <ul style="list-style-type: none"> • CASE_IGNORE Fold everything to lowercase for matching • CASE_INSENSITIVE Fold only tokens with initial caps to lowercase • CASE_FOLD_DIGITS Fold all (and only) tokens with a digit • CASE_SENSITIVE Perform no case folding |
| Stemmer | <ul style="list-style-type: none"> • BIOLEMMATIZER A stemmer specific for biomedical literature • PORTER A stemmer that removes the commoner morphological and inflexional endings from words in English • NONE No word stemming |
| StopWords | <ul style="list-style-type: none"> • PUBMED A list of stop words obtained analyzing Pubmed papers • NONE No stop words |
| OrderIndependentLookup | <ul style="list-style-type: none"> • ON Ordering within span is ignored (i.e. 'Breast cancer' would equal 'Cancer breast') • OFF Ordering is taken into consideration |
| FindAllMatches | <ul style="list-style-type: none"> • YES All the matches within the span are found |

• NO Only the longest match within the span will be returned
 • EXACT_ONLY Only exact synonyms are considered
 • ALL All synonym types are included

paramValueIndex
 An integer value to index the 576 parameter combinations

Value

instance of the class COptions set to the default combination of parameters

Examples

```
op <- COptions()
```

COptions-class	<i>Class to set the options to run the EntityFinder</i>
----------------	---

Description

COptions is a class that represents Conceptmapper configurations. It allows users to set the possible combinations of different parameters for Conceptmapper running.

This method shows the list of options to run the Entity finder

Usage

```
## S4 method for signature 'COptions'
show(object)
```

Arguments

object COptions instance

Details

The following methods can be applied to COptions

```

show
paramValueIndex
paramValueIndex<-
SearchStrategy
SearchStrategy<-
CaseMatch
CaseMatch<-
Stemmer
Stemmer<-
StopWords
StopWords<-
OrderIndependentLookup
OrderIndependentLookup
FindAllMatches
FindAllMatches<-
SynonymType
SynonymType<-

```

Value

the list of options

Slots

paramValueIndex An integer value to index the 576 parameter combinations

SearchStrategy The matching strategy for finding concepts in the input text

- CONTIGUOUS_MATCH Longest match of contiguous tokens within enclosing span
- SKIP_ANY_MATCH Longest match of not-necessarily contiguous tokens
- SKIP_ANY_MATCH_ALLOW_OVERLAP Longest match of not-necessarily contiguous tokens, overlapping matches are allowed

CaseMatch • CASE_IGNORE Fold everything to lowercase for matching

- CASE_INSENSITIVE Fold only tokens with initial caps to lowercase
- CASE_FOLD_DIGITS Fold all (and only) tokens with a digit
- CASE_SENSITIVE Perform no case folding

Stemmer • BIOLEMMATIZER A stemmer specific for biomedical literature

- PORTER A stemmer that removes the commoner morphological and inflexional endings from words in English
- NONE No word stemming

StopWords • PUBMED A list of stop words obtained analyzing Pubmed papers

- NONE No stop words

OrderIndependentLookup • ON Ordering within span is ignored (i.e. 'Breast cancer' would equal 'Cancer breast')

- OFF Ordering is taken into consideration

FindAllMatches • YES All the matches within the span are found

- NO Only the longest match within the span will be returned

SynonymType • EXACT_ONLY Only exact synonyms are considered

- ALL All synonym types are included

Examples

```
options <- new('COptions')
opt <- COptions()
show(opt)
```

collapse

collapse

Description

This method collapses semantic states in an Onassis object.

Usage

```
collapse(onassis = NA, simil_thresh)
```

```
## S4 method for signature 'Onassis'
collapse(onassis = NA, simil_thresh)
```

Arguments

onassis instance of class [Onassis-class](#)
 simil_thresh the semantic similarity threshold to use to merge similar semantic sets

Value

a new object of class [Onassis-class](#) with collapsed annotations for the entities and a new similarity matrix of similarities between newly created semantic sets

Examples

```
geo_chip <- readRDS(system.file('extdata', 'vignette_data',
'GEO_human_chip.rds', package='Onassis'))
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
onassis_results <- annotate(geo_chip, 'OBO', dictionary=obo)
entities <- entities(onassis_results)
entities(onassis_results) <- entities[sample(nrow(entities), 15),]
onassis_results <- sim(onassis_results)
collapsed_onassis <- collapse(onassis_results, 0.9)
```

 compare

 compare

Description

This method compares lists of scores (e.g. gene expression values, gene copy numbers, binding factors intensity) associated to the annotated entities (samples) of an [Onassis](#) object according to the semantic sets obtained from the annotation step. For [Onassis](#) objects annotated with a single ontology the method applies a test function to determine differences between subsets of the scores in one semantic set compared to subsets of scores in any other semantic set. For [Onassis](#) objects containing annotated entities with two ontologies, the first ensemble of semantic sets (e.g. cell lines) will be used as the main container to organize samples and the comparisons will be carried out between the entities belonging to different semantic sets of the second ontology (e.g. disease) within each semantic set defined from the first ontology.

Usage

```
compare(onassis, ...)  
  
## S4 method for signature 'Onassis'  
compare(onassis, score_matrix = NA, by = "row",  
      fun_name = "wilcox.test", fun_args = list(), padj = FALSE)
```

Arguments

onassis instance of class [Onassis-class](#)
 ... Optional parameters
 score_matrix a matrix of scores containing on the rows genomic units and on the columns the samples annotated in the entities
 by 'row' if the test refers to single genomic units in multiple conditions, 'col' if the test compares all the genomic units across different conditions. Defaults to row

fun_name	name of the test to apply
fun_args	list of arguments needed by the function
padj	TRUE if multiple test correction is needed

Details

The entities slot of an Onassis object can contain annotations with concepts from one ontology or two ontologies. The function compare separates these two scenarios.

ONE ONTOLOGY CASE: The entities (samples) are assigned to their corresponding semantic set (e.g. cell lines) A score matrix should have as rows genomic units (gene names, genomic regions, mutation identifiers...) and as columns the identifiers of all the entities annotated and belonging to semantic sets found in the sample_id field. The score for each entry of the matrix can be any biological measurement (gene expression RPKMs, peak intensity, copy numbers...) Importantly, each semantic set can contain a different number of annotated entities (samples) and thus a single vector of scores (in case only 1 entity belongs to 1 semantic set) or a subset of the columns of the scores matrix If by is "row" then the function provided as fun_name will be applied to compare, genomic unit by genomic unit (rows of the score matrix), all the possible couples of semantic sets. The function can be any multiple statistical test function taking as parameters: - two numeric vectors of potentially different lengths (for genomic unit in row i, n samples in semantic set 1 and m samples in semantic set 2) - other optional arguments needed by the function passed as a list in fun_args The function should always provide as result a "statistic" and "p.value" In case the padj is set to TRUE the bonferroni correction will be applied for multiple tests If by is "col" then the function provided as fun_name will be applied to the couples of semantic sets considering all the possible values for the genomic units. In this case the test function should take as parameters - two matrices with potentially different number of columns - other optional arguments needed by the function can be passed as fun_args **TWO ONTOLOGIES CASE:** The comparisons in this case will be carried out considering for each semantic set defined from the primary ontology, all the possible couples of semantic sets generated from the second ontology (e.g. for each cell line, different diseases) within the first ontology semantic sets

Value

The results of the comparison between semantic classes. In case of one ontology a square matrix whose rows and columns contain the semantic sets. For each couple of semantic sets i and j if by is 'row' the element in the row i and column j is a data frame with biological entities (genes, regions..) and for each entity, the results of the statistics function and p.value. For multiple test functions if padj = TRUE then also the Bonferroni correction will be reported. For each couple of semantic sets i and j if by is 'col' the element at row i and column j is a couple of values with the result of the statistic and corresponding p.value In case of two ontologies the result will be a list named with the first level semantic sets and each element of the list will be the result of the comparisons between second level semantic sets within first level semantic sets. Depending on the by parameter, the elements of the list will be matrices or vectors as for the one ontology case.

Examples

```
#Loading ChIP-seq data
geo_chip <- readRDS(system.file('extdata', 'vignette_data', 'GEO_human_chip.rds', package='Onassis'))
#Sampling 30 samples
geo_chip <- geo_chip[sample(nrow(geo_chip), 30) ,]
# Loading the obo ontology for cell lines
obo1 <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
# Loading the obo ontology for diseases
obo2 <- system.file('extdata', 'sample.do.obo', package='OnassisJavaLibs')
```

```

#Annotating cell lines
onassis_results1 <- annotate(geo_chip, 'OBO', dictionary=obo1)
#Annotating diseases
onassis_results2 <- annotate(geo_chip, 'OBO', dictionary=obo2)
# Creating a score matrix
n <- length(unique(geo_chip$sample_accession))
m <- 50
score_matrix <- matrix(sample(0:1, m * n, replace = TRUE), m, n)
colnames(score_matrix) <- unique(geo_chip$sample_accession)
rownames(score_matrix) <- paste0('gene_', seq(1, m, 1))
# Merging the annotations from the two ontologies in a single object
my_onassis <- mergeonassis(onassis1 = onassis_results1, onassis2 = onassis_results2)
scores(my_onassis) <- score_matrix
# Comparing the scores associated to samples belonging to cell line semantic sets
# By default the wilcox.test will be applied to each of the 50 genes
scores(onassis_results1) <- score_matrix
gene_by_gene_cell_differences <- compare(onassis_results1)
head(gene_by_gene_cell_differences[1,2])
# Applying the same wilcox.test but obtaining a multiple test correction
gene_by_gene_cell_differences_ADJ <- compare(onassis_results1, padj=TRUE)
head(gene_by_gene_cell_differences_ADJ[1,2])
# Comparing disease genes across all the tissue semantic sets with wilcox.test and applying Bonferroni correction
gene_by_gene_disease_differences <- compare(my_onassis, by='row', padj=TRUE, fun_name='wilcox.test')
# Comparing diseases in cell line semantic sets
disease_differences <- compare(my_onassis, by='col', fun_name='wilcox.test')
# Using a personalized function
mykruskal <- function(x, y, params){
  kruskal.test(as.list(c(x, y)))}
cell_differences_personalized <- compare(onassis_results1, by='col', fun_name='mykruskal')

```

connectToGEODB

connectToGEODB

Description

This method allows users to connect to the GEOmetadb downloaded. If no parameter is provided than the function retrieves the database in sqlite format and returns a connection to query the database

Usage

```
connectToGEODB(sqliteFilePath = NULL, download = FALSE,
  destdir = getwd())
```

Arguments

sqliteFilePath optional SQLite full file path of the SQLite database if already downloaded
download If TRUE allow the automatic downloading of the database file.
destdir optional destination directory. Current working directory is the default

Value

A connection to the GEOmetadb

Examples

```
## Not run:
geo_connection <- connectToGEODB(download=TRUE)

## End(Not run)
if(file.exists('GEOmetadb.sqlite')){
  geo_con <- connectToGEODB()
} else {
  message('Please provide GEOmetadb.sqlite file')
}
```

create_score_matrix createScoreMatrix

Description

This functions allows the creation of a score matrix for genomic regions overlapping the genomic regions provided as reference (e.g. promoter regions). Each entry of the matrix corresponds to 1 if the sample represented on the column overlaps the genomic interval represented on the row.

Usage

```
create_score_matrix(ref_granges, granges_list)
```

Arguments

`ref_granges` An object of type GRanges to be considered as the reference GRanges
`granges_list` A list of GRanges, one for each sample, to be mapped on the reference granges

Value

A logical score matrix where the number of rows corresponds to the number of different genomic intervals in the `ref_granges` object and each column is associated to a Grange in the `granges_list`. `Score(i, j)` is set to 1 if the `j`-th sample has a interval overlapping the one in the `i`-th row of the reference GRanges.

Examples

```
granges <- readRDS(system.file('extdata', 'sample_granges.rds', package='Onassis'))
ref_granges <- granges[[1]]

for(i in 2:length(granges)) {
  ref_granges <- GenomicRanges::union(ref_granges, granges[[i]])
}
score_mat <- create_score_matrix(ref_granges, granges)
```

dictInfo	<i>Get and Set the dicInfo slot for the CMdictionary class</i>
----------	--

Description

Method to get and set a list of info about the dictionary

Usage

```
dictInfo(.Object)

dictInfo(.Object) <- value

## S4 method for signature 'CMdictionary'
dictInfo(.Object)

## S4 replacement method for signature 'CMdictionary'
dictInfo(.Object) <- value
```

Arguments

.Object	An instance of class CMdictionary
value	list of details about the dictionary

Value

list of details about the dictionary
 object of class [CMdictionary](#)

Examples

```
dictionary <- new('CMdictionary')
dictInfo(dictionary)
dictionary <- new('CMdictionary')
dictInfo(dictionary) <-
list(Dictionary_type = 'ENTREZ from OrgDb', Dictionary_source = 'OrgDb')
```

dictionary	dictionary<-
------------	--------------

Description

Method to get and set the dictionary slot of the class [Onassis-class](#)

Usage

```
dictionary(object)

dictionary(object) <- value

## S4 replacement method for signature 'Onassis'
dictionary(object) <- value

## S4 method for signature 'Onassis'
dictionary(object)
```

Arguments

object	instance of class Onassis-class
value	the path of an OBO file

Value

The path of the dictionary in case of get, the instance of Onassis with new dictionary in case of set

Examples

```
onassis <- Onassis()
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
dictionary(onassis) <- obo
o <- Onassis()
dictionary(o)
```

dictRef

dictRef

Description

This method retrieves and sets the java reference the conceptmapper dictionary

Usage

```
dictRef(.Object)

dictRef(.Object) <- value

## S4 method for signature 'CMdictionary'
dictRef(.Object)

## S4 replacement method for signature 'CMdictionary'
dictRef(.Object) <- value
```

Arguments

.Object	An instance of class CMdictionary
value	the reference of a XML Conceptmapper dictionary file already created

Value

java reference to the Conceptmapper dictionary

Examples

```
dictionary <- new('CMdictionary')
dictRef(dictionary)
dictionary <- new('CMdictionary')
dict_file <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
dictRef(dictionary) <- .jnew('java/io/File', dict_file)
```

dictTypes

dictTypes

Description

A function to show the pre-defined Conceptmapper dictionary types wwith their descriptions

Usage

```
dictTypes()
```

Value

list of types that can be used to create a Conceptmapper dictionary

Examples

```
dictTypes()
```

dict_location

dict_location

Description

A method to get and set the location of [CMdictionary](#)

Usage

```
dict_location(.Object)
```

```
dict_location(.Object) <- value
```

```
## S4 method for signature 'CMdictionary'
dict_location(.Object)
```

```
## S4 replacement method for signature 'CMdictionary'
dict_location(.Object) <- value
```

Arguments

.Object	An instance of class CMdictionary
value	the path of the new dictionary

Value

the path of the dictionary in the file system

Examples

```
dictionary <- new('CMdictionary')
loc <- dict_location(dictionary)
dictionary <- new('CMdictionary')
dict_location(dictionary) <- getwd()
```

entities	entities<-
----------	------------

Description

Method to get and set the entities slot of the class [Onassis-class](#)

Usage

```
entities(object)
```

```
entities(object) <- value
```

```
## S4 replacement method for signature 'Onassis'
entities(object) <- value
```

```
## S4 method for signature 'Onassis'
entities(object)
```

Arguments

object	instance of class Onassis-class
value	a data frame with annotated entities

Value

the entities of the Onassis object in case of get and the Onassis object with new entities in case of set

Examples

```
onassis <- Onassis()
entities(onassis) <- data.frame()
```

```
o <- Onassis()
entities(o)
```

EntityFinder	EntityFinder
--------------	--------------

Description

this function creates instances of the class [EntityFinder](#)

Usage

```
EntityFinder(input, dictionary, options = NA, outDir = tempdir(),
             multipleDocs = FALSE)
```

Arguments

input	the file, directory, or data frame with the text to annotate
dictionary	A dictionary of the type CMdictionary or the path to an already created Conceptmapper XML file
options	an object of class CMoptions . If NA, the default configuration will be set.
outDir	the directory where the annotated files will be stored
multipleDocs	TRUE when multiple documents are loaded from a single file with each row representing a document. The file should have two columns. The first for the unique document identifier and the second for the textual descriptions

Value

dataframe of annotations

Examples

```
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
sample_dict <- CMdictionary(input=obo, outputDir=getwd(), synonymType='ALL')
myopts <- CMoptions()
paramValueIndex(myopts) <- 40
entities <- EntityFinder(input=readRDS(system.file('extdata', 'vignette_data', 'GEO_human_chip.rds', package='OnassisJavaLibs')))
```

EntityFinder-class	<i>EntityFinder class to create a Conceptmapper instance</i>
--------------------	--

Description

EntityFinder is a class that wraps a Conceptmapper pipeline using the CCP UIMA Type System <https://github.com/UCDenver-ccp/ccp-nlp>. The pipeline includes a sentence detector, offset tokenizer and retrieves concepts from dictionaries built from OBO/OWL formatted ontology files.

Details

The following methods can be applied to EntityFinder

[findEntities](#)
[annotatedDF](#)

The methods can be automatically called using the function [EntityFinder](#)

Slots

typeSystemRef The reference to the Java object representing the type system

Examples

```
finder <- new('EntityFinder')
```

experiment_types	experiment_types
------------------	------------------

Description

This method retrieves the experiment types stored in GEOmetadb

Usage

```
experiment_types(GEOcon)
```

Arguments

GEOcon connection to the SQLite GEOmetadb database

Value

A character vector with all the possible experiment values

Examples

```
if(file.exists('GEOmetadb.sqlite')){
  geo_con <- connectToGEODB('GEOmetadb.sqlite')
  experiments <- experiment_types(geo_con)
}else{
  print('You need to download GEOmetadb.sqlite to run this example')
}
```

filterconcepts	filterconcepts
----------------	----------------

Description

This method filters unwanted concepts from the entities of an [Onassis-class](#) object

Usage

```
filterconcepts(onassis, concepts_to_filter)
```

```
## S4 method for signature 'Onassis'
filterconcepts(onassis, concepts_to_filter = c())
```

Arguments

onassis An object of class `Onassis-class` with already annotated entities
concepts_to_filter A vector with unwanted concepts

Value

the instance of `Onassis-class` with filtered entities

Examples

```
geo_chip <- readRDS(system.file('extdata', 'vignette_data',  
'GEO_human_chip.rds', package='Onassis'))  
geo_chip <- geo_chip[sample(nrow(geo_chip), 15) ,]  
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')  
onassis_results <- annotate(geo_chip, 'OBO', dictionary=obo)  
filtered_onassis <- filterconcepts(onassis_results, c('cell'))
```

filterTerms

filterTerms

Description

filterTerms allows users to remove a set of defined terms from the results of the annotation provided by Onassis

Usage

```
filterTerms(annotated_df, termlist = c())
```

Arguments

annotated_df a data frame resulting from the annotation process of Onassis
termlist the list of terms to be filtered out from the annotations

Value

a data frame with removed annotations

Examples

```
metadatafile <- readRDS(system.file('extdata', 'vignette_data',  
'GEO_human_chip.rds', package='Onassis'))  
healthy_gsms <- findHealthy(metadatafile)
```

FindAllMatches *Method FindAllMatches*

Description

Method to get and set the FindAllMatches parameter

Usage

```
FindAllMatches(x)
```

```
FindAllMatches(x) <- value
```

```
## S4 method for signature 'COptions'  
FindAllMatches(x)
```

```
## S4 replacement method for signature 'COptions'  
FindAllMatches(x) <- value
```

Arguments

x	instance of class COptions
value	<ul style="list-style-type: none">• YES All the matches within the span are found• NO Only the longest match within the span will be returned

Value

The FindAllMatches corresponding to the current options when used as get, the new COptions object with updated parameters when used to set.

Examples

```
opts <- COptions()  
FindAllMatches(opts)  
opts <- COptions()  
FindAllMatches(opts) <- 'YES'
```

findEntities *Method findEntities*

Description

This method finds concepts from a Conceptmapper Dictionary of type [CMdictionary](#) in a given directory or in a single pipe separated file containing a named document in each row, with a specified configuration of type [COptions](#). This is a method of the [EntityFinder-class](#)

Usage

```
findEntities(object, inputDirOrFile, multipleDocs = FALSE,
             outDir = tempdir(), configOpt, cmDict)

## S4 method for signature
## 'EntityFinder,character,logical,character,CMoptions,CMdictionary'
findEntities(object,
             inputDirOrFile, multipleDocs = FALSE, outDir = tempdir(), configOpt,
             cmDict)
```

Arguments

object	instance of the class EntityFinder
inputDirOrFile	the directory where the files to annotate are stored or the text file to annotate. A single file containing in each row sample names, the symbol and the description of the sample is also allowed.
multipleDocs	TRUE if a single file containing different text sources has been given as inputDirOrFile. FALSE if each text is in a separate file. Defaults to FALSE
outDir	The directory where the Conceptmapper annotated files are stored. Default: the system tmp directory.
configOpt	Object of type CMoptions in which the parameters to run Conceptmapper are stored
cmDict	Instance of class CMdictionary or the file path of an already created CMdictionary

Value

A data frame of annotations containing the sample name, the id of the OBO concept, the corresponding name, the part of the text containing the annotation

Examples

```
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
dict <- CMdictionary(inputFileOrDb=obo, outputDir=getwd(), synonymType='ALL')

opts <- CMoptions()
ef <- new('EntityFinder')
annotations <- findEntities(ef,
system.file('extdata', 'test_samples', 'test_samples.txt', package='Onassis'), multipleDocs=TRUE, outDir=getwd(),
configOpt=opts, cmDict=dict)
```

findHealthy

findHealthy

Description

findHealthy annotates as 'Healthy' the samples whose metadata matches with one of the elements of a list of sentence used to describe the normal disease state or healthy.

Usage

```
findHealthy(metadata_df)
```

Arguments

`metadata_df` a data frame where the first column corresponds to the identifier of a sample # and the other columns the metadata relative to the sample

Value

a data frame with healthy samples annotated as 'Healthy'

Examples

```
metadatafile <- readRDS(system.file('extdata', 'vignette_data',
  'GEO_human_chip.rds', package='Onassis'))
```

```
healthy_gsms <- findHealthy(metadatafile)
```

```
getGEOMetadata      getGEOMetadata
```

Description

This method retrieves the descriptive fields of the samples in GEO for a given `experiment_type`, `organism` or `platform`.

Usage

```
getGEOMetadata(geo_con, experiment_type = NA, organism = NA,
  gpl = NA)
```

Arguments

`geo_con` connection to the SQLite GEOmetadb database

`experiment_type` The type of experiment. Allowed values can be obtained through the function `experiment_types`

`organism` Optional type of organism. Allowed species can be obtained using the function `organism_types`. If no organism is passed as parameter the query will retrieve all the organisms

`gpl` Optional platform identifier in case a platform based query has to be executed

Value

A data frame with the queried samples' metadata

Examples

```

if(file.exists('GEOmetadb.sqlite')){
  geo_con <- connectToGEODB('GEOmetadb.sqlite')
  methylation <- getGEOMetadata(geo_con,
    'Methylation profiling by high throughput sequencing', 'Homo sapiens')
  expression <- getGEOMetadata(geo_con,
    'Expression profiling by array', 'Homo sapiens', 'GPL570')
}else{
  print('You need to download GEOmetadb.sqlite to run this example')
}

```

groupConfig

groupConfig

Description

This method shows the value of the groupwise configuration used to compute semantic similarities between groups of concepts.

Sets the groupwise measure to compute the semantic similarity between groups of concepts. For available measures use the method `listSimilarities()`.

Usage

```

groupConfig(object)

groupConfig(object) <- value

## S4 method for signature 'Similarity'
groupConfig(object)

## S4 replacement method for signature 'Similarity'
groupConfig(object) <- value

```

Arguments

object	instance of class Similarity-class
value	See details

Details

The following measures are indirect groupwise measures, meaning that they are used to aggregate individual pairwise measures.

- 'min': Minimum of the pairwise similarities of the concepts in the two groups
- 'average': Average of the pairwise similarities of the concepts in the two groups
- 'max': Max of the pairwise similarities of the concepts in the two groups
- 'bma': Best match average
- 'bmm': Best match max

Direct groupwise measures directly compare the sets of concepts considering the features of both sets.

- 'ui': Considers the intersection and the union of the set of ancestors of the two groups of concepts:

$$sim(group_u, group_v) = |intersection(A(group_u), A(group_v))| / |union(A(group_u), A(group_v))|$$

- 'nto_max': Normalized max Term Overlap, computes the groupwise semantic similarity considering the inclusive set of ancestors of the two groups of concepts.

$$sim(group_u, group_v) = |intersection(A(group_u), A(group_v))| / max(|A(group_u)|, |A(group_v)|)$$

- 'lee': Computes the groupwise semantic similarity considering the inclusive set of ancestors of the two groups of concepts.

$$sim(group_u, group_v) = |union(A(group_u), A(group_v))|$$

- 'lp': Computes the groupwise semantic similarity between two groups of concepts as the depth of the longest shared path from the root node

- 'gic': Computes the groupwise semantic similarity between two groups of concepts as the ration between the information content of the concepts in the intersection of the ancestors in the two groups and the information content of the concepts in the union of the ancestors in the two groups.

$$sim(group_u, group_v) = IC_{intersection} / IC_{union}$$

- 'batet': Computes the groupwise semantic similarity between two groups of concepts considering the union and intersection of ancestors normalized on the number of concepts in the ontology.

$$sim(group_u, group_v) = |(union(A(group_u), A(group_v)) - intersection(A(group_u), A(group_v)))| / (|union_{tot_concepts}|)$$

Value

groupwise configured measure for the similarity object provided as input instance of the Similarity class with the new groupwise option.

Examples

```
sim <- new('Similarity')
groupConfig(sim)
sim <- new('Similarity')
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
ontology(sim) <- obo
groupConfig(sim) <- 'ui'
```

groupsim

Method groupsim

Description

This method computes the semantic similarity between two groups of terms of a given ontology.

Usage

```
groupsim(object, termList1, termList2)
```

```
## S4 method for signature 'Similarity,character,character'
groupsim(object, termList1,
          termList2)
```

Arguments

object	instance of class Similarity-class
termList1	A vector of URIs of ontology terms in the format http://purl.obolibrary.org/obo/Ontology_id (e.g http://purl.obolibrary.org/obo/BTO_0004732)
termList2	A vector of URIs of ontology terms

Value

the semantic similarity of the two provided groups of concepts

Examples

```
sim <- new('Similarity')
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
ontology(sim) <- obo
pairwiseConfig(sim) <- listSimilarities()$pairwiseMeasures[9]
groupConfig(sim) <- listSimilarities()$groupwiseMeasures[3]
similarity <- groupsim(sim, c('http://purl.obolibrary.org/obo/CL_0000542',
  'http://purl.obolibrary.org/obo/CL_0000236'),
  c('http://purl.obolibrary.org/obo/CL_0000000'))
similarity
```

groupwiseConfigRef	groupwiseConfigRef
--------------------	--------------------

Description

This method shows the value of the groupwise configuration used to compute semantic similarities between groups of concepts.

Sets the groupwise measure to the reference of a groupwise measure to the semantic similarity between groups of concepts. For available measures see the `groupConfig` function's details.

Usage

```
groupwiseConfigRef(object)

groupwiseConfigRef(object) <- value

## S4 method for signature 'Similarity'
groupwiseConfigRef(object)

## S4 replacement method for signature 'Similarity'
groupwiseConfigRef(object) <- value
```

Arguments

object	instance of class Similarity-class
value	one of the groupwise measures. See groupConfig

Value

groupwise configured measure for the similarity object provided as input instance of the Similarity class with the new groupwise option.

Examples

```
sim <- new('Similarity')
groupwiseConfigRef(sim)
sim <- new('Similarity')
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
groupwiseConfigRef(sim) <- 'ui'
```

 icConfig

 icConfig

Description

This method retrieves the configuration of the intrinsic information content measure

This method sets the configuration of the intrinsic information content measure by taking as parameter the short flag associated to the information measure. To have details about the available short flags see the [pairwiseConfig](#) help

Usage

```
icConfig(object)

icConfig(object) <- value

## S4 method for signature 'Similarity'
icConfig(object)

## S4 replacement method for signature 'Similarity'
icConfig(object) <- value
```

Arguments

object	instance of class Similarity-class
value	the information content measure selected.

Value

The measure used to compute concepts' information content

The similarity object with the new information content measure set

Examples

```
sim <- new('Similarity')
icConfig(sim)
sim <- new('Similarity')
icConfig(sim) <- 'sanchez'
```

listCMOptions
listCMOptions

Description

This method retrieves all the possible parameters combinations for Conceptmapper.

paramValueIndex An integer value to index the 576 parameter combinations

SearchStrategy The matching strategy for finding concepts in the input text

- CONTIGUOUS_MATCH Longest match of contiguous tokens within enclosing span
- SKIP_ANY_MATCH Longest match of not-necessarily contiguous tokens
- SKIP_ANY_MATCH_ALLOW_OVERLAP Longest match of not-necessarily contiguous tokens, overlapping matches are allowed

CaseMatch • CASE_IGNORE Fold everything to lowercase for matching

- CASE_INSENSITIVE Fold only tokens with initial caps to lowercase
- CASE_FOLD_DIGITS Fold all (and only) tokens with a digit
- CASE_SENSITIVE Perform no case folding

Stemmer • BIOLEMMATIZER A stemmer specific for biomedical literature

- PORTER A stemmer that removes the commoner morphological and inflexional endings from words in English
- NONE No word stemming

StopWords • PUBMED A list of stop words obtained analyzing Pubmed papers

- NONE No stop words

OrderIndependentLookup • ON Ordering within span is ignored (i.e. 'Breast cancer' would equal 'Cancer breast')

- OFF Ordering is taken into consideration

FindAllMatches • YES All the matches within the span are found

- NO Only the longest match within the span will be returned

SynonymType • EXACT_ONLY Only exact synonyms are considered

- ALL All synonym types are included

Usage

```
listCMOptions()
```

Value

The data frame with all the possible parameter combinations

Examples

```
o <- listCMOptions()
```

listSimilarities	listSimilarities
------------------	------------------

Description

This method shows a list of the possible measures to compute pairwise and groupwise semantic similarity between concepts

Usage

```
listSimilarities()
```

Value

the list of pairwise, information content and groupwise measures to compute the semantic similarities

Examples

```
s <- listSimilarities()
```

mergeonassis	mergeonassis
--------------	--------------

Description

This method unifies the entities of two Onassis objects

Usage

```
mergeonassis(onassis1 = NA, onassis2 = NA)
```

```
## S4 method for signature 'Onassis,Onassis'  
mergeonassis(onassis1 = NA, onassis2 = NA)
```

Arguments

onassis1 instance of class [Onassis-class](#)

onassis2 instance of class [Onassis-class](#)

Value

new object of type [Onassis-class](#) with merged entities

Examples

```

geo_chip <- readRDS(system.file('extdata', 'vignette_data',
'GEO_human_chip.rds', package='Onassis'))
geo_chip <- geo_chip[sample(nrow(geo_chip), 15) ,]
obo1 <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
obo2 <- system.file('extdata', 'sample.do.obo', package='OnassisJavaLibs')
onassis_results1 <- annotate(geo_chip, 'OBO', dictionary=obo1)
onassis_results2 <- annotate(geo_chip, 'OBO', dictionary=obo2)
onassis_results <- mergeonassis(onassis_results1, onassis_results2)

```

multisim

multisim

Description

This method computes the semantic similarity between samples annotated with different ontology terms from different ontologies

Usage

```

multisim(similarities, annotations, sample1, sample2, aggregating_function)

## S4 method for signature 'list,list,character,character'
multisim(similarities,
  annotations, sample1, sample2, aggregating_function = "mean")

```

Arguments

similarities	a list of Similarity instances, one for each ontology used to annotate the data
annotations	a list of annotated data frames obtained using <code>annotateDF</code> or <code>findEntities</code> , one for each ontology
sample1	the name of a sample in annotations
sample2	the name of a sample in annotations
aggregating_function	A function used to aggregate the single similarities obtained from each ontology annotation. The function should be applied to a numeric vector. The default value is 'mean'

Value

The aggregate semantic similarity between the samples `sample1` and `sample2`

Examples

```

ef <- new('EntityFinder')

opts <- COptions()
cell_dict_file <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
sample_dict <- CMdictionary(cell_dict_file, outputDir=getwd(),
synonymType='ALL')

```

```

samples <- findEntities(ef, system.file('extdata', 'test_samples',
'test_samples.txt',
package='Onassis'), outDir=getwd(), multipleDocs=TRUE, configOpt=opts,
  cmDict=sample_dict)
d_dict_file <- system.file('extdata', 'sample.do.obo', package='OnassisJavaLibs')
disease_dict <- CMdictionary(d_dict_file, outputDir=getwd(), synonymType='ALL')
disease <- findEntities(ef, system.file('extdata', 'test_samples',
'test_samples.txt', package='Onassis'),
  multipleDocs=TRUE, outDir=getwd(), configOpt=opts,
  cmDict=disease_dict)

cell_sim <- new('Similarity')
ontology(cell_sim) <- cell_dict_file

disease_sim <- new('Similarity')
ontology(disease_sim) <- d_dict_file

pairwiseConfig(cell_sim) <- listSimilarities()$pairwiseMeasures[9]
pairwiseConfig(disease_sim) <- listSimilarities()$pairwiseMeasures[9]
groupConfig(cell_sim) <- listSimilarities()$groupwiseMeasures[3]
groupConfig(disease_sim) <- listSimilarities()$groupwiseMeasures[3]
similarity <- multisim(list(cell_sim, disease_sim),
  list(samples, disease),
  as.character(as.vector(samples[1,1])),
  as.character(as.vector(samples[5,1])), 'mean')

```

OnASSiS

OnASSiS (Ontology Annotations and Semantic Similarity software)

Description

OnASSiS (Ontology Annotations and Semantic Similarity software) is a package for the annotation of any given text with concepts from biomedical ontologies that also provides features to relate the concepts using semantic similarity metrics.

This constructor instantiates an Onassis object.

Usage

```

Onassis(dictionary = NA_character_, entities = data.frame(),
  similarity = matrix(), scores = matrix())

```

Arguments

dictionary	The path of the dictionary file
entities	a data frame to store entities
similarity	A matrix of the similarities between entities
scores	The result of comparisons of the elements in the entities

Details

OnASSiS package

OnASSiS (Ontology Annotations and Semantic Similarity software) is a package that uses Conceptmapper, an Apache UIMA (Unstructured Information Management Architecture) <https://uima.apache.org/downloads/sandbox/ConceptMapperAnnotatorUserGuide/ConceptMapperAnnotatorUserGuide.html> dictionary lookup tool to retrieve dictionary terms in a given text.

In particular a Conceptmapper wrapper specific for the biomedical domain, ccp-nlp, (<https://github.com/UCDenver-ccp/ccp-nlp>) has been personalized to retrieve concepts from OBO ontologies in a given text with different options.

The package also provides the possibility to annotate Gene Expression Omnibus (GEO) metadata for stored experiments and samples.

Different annotated sets of text can be then compared using semantic similarity metrics based on the structure of the biomedical ontologies. The semantic similarity module has been obtained using the Java slib (<http://www.semantic-measures-library.org/sml/>)

Value

An object of type Onassis that can be used to analyze metadata

Examples

```
onassis <- Onassis()
```

Onassis-class

Onassis-class

Description

Onassis is a container class for annotating samples metadata with concepts from dictionaries/ontologies, creating semantic sets of unique annotations, computing the distances between different semantic sets and eventually comparing the different identified conditions.

Details

The following methods can be applied to Onassis

```
annotate
collapse
compare
dictionary
simil
entities
scores
sim
```

Slots

dictionary One or more input dictionaries to annotate samples metadata
 entities a data frame containing the result of the annotation of the input with ontology terms
 similarity A matrix of the similarities between the entries in the entities slot
 scores An optional score matrix containing genomic units on the rows (genes, regions) and on the columns the elements on the rows of the entities slot

ontology	ontology<-
----------	------------

Description

This method creates a semantic graph to compute semantic similarity between concepts. It takes as input an OBO ontology in RDF, OWL or OBO format.

This method shows the ontology.

Usage

```
ontology(object)

ontology(object) <- value

## S4 replacement method for signature 'Similarity'
ontology(object) <- value

## S4 method for signature 'Similarity'
ontology(object)
```

Arguments

object	instance of class Similarity-class
value	The path of an ontology file

Value

The Similarity object where 'ontology' slot refers to the Java graph created
 Ontology object

Examples

```
sim <- new('Similarity')
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
ontology(sim) <- obo
sim <- new('Similarity')
ontology(sim)
```

```
OrderIndependentLookup
      OrderIndependentLookup
```

Description

Method to get and set the OrderIndependentLookup parameter

Usage

```
OrderIndependentLookup(x)

OrderIndependentLookup(x) <- value

## S4 method for signature 'COptions'
OrderIndependentLookup(x)

## S4 replacement method for signature 'COptions'
OrderIndependentLookup(x) <- value
```

Arguments

x	instance of class COptions
value	<ul style="list-style-type: none"> • ON Ordering within span is ignored (i.e. 'Breast cancer' would equal 'Cancer breast') • OFF Ordering is taken into consideration

Value

The OrderIndependentLookup corresponding to the current options when used as get, the new COptions object with updated parameters when used to set.

Examples

```
opts <- COptions()
OrderIndependentLookup(opts)

opts <- COptions()
OrderIndependentLookup(opts) <- 'ON'
```

```
organism_types      organism_types
```

Description

This method retrieves the allowed organisms in GEOmetadb

Usage

```
organism_types(geo_con)
```

Arguments

geo_con connection to the SQLite GEOmetadb database

Value

A character vector with all the possible organism values

Examples

```
if(file.exists('GEOmetadb.sqlite')){
  geo_con <- connectToGEODB('GEOmetadb.sqlite')
  species <- organism_types(geo_con)
}else{
  print('You need to download GEOmetadb.sqlite to run this example')
}
```

pairsim

pairsim

Description

This method computes the semantic similarity between two terms of a given ontology.

Usage

```
pairsim(object, term1, term2)
```

```
## S4 method for signature 'Similarity,character,character'
pairsim(object, term1, term2)
```

Arguments

object instance of class [Similarity-class](#)

term1 The URI of the ontology term in the format http://purl.obolibrary.org/obo/Ontology_id (e.g 'http://purl.obolibrary.org/obo/CL_0000542')

term2 The URI of the ontology term

Value

the semantic similarity of the two provided concepts

Examples

```
sim <- new('Similarity')
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
ontology(sim) <- obo
pairwiseConfig(sim) <- listSimilarities()$pairwiseMeasures[9]
similarity <- pairsim(sim, 'http://purl.obolibrary.org/obo/CL_0000542',
  'http://purl.obolibrary.org/obo/CL_0000236')
```

pairwiseConfig	pairwiseConfig
----------------	----------------

Description

This method shows the value of the pairwise configuration.

and configures the pairwise measure to compute semantic similarity between two concepts of a given ontology. To set the pairwise measure one of the available short flags described in details should be used.

Usage

```
pairwiseConfig(object)

pairwiseConfig(object) <- value

## S4 method for signature 'Similarity'
pairwiseConfig(object)

## S4 replacement method for signature 'Similarity'
pairwiseConfig(object) <- value
```

Arguments

object	instance of class Similarity-class
value	See details

Details

The following measures can be used to compute semantic similarities between two concepts.

- 'edge_rada_lca': Computes the similarity of two concepts based on the shortest path linking the two concepts.

$$sim(u, v) = 1/sp(u, v)$$
- 'edge_wupalmer': Computes the similarity of two concepts based on the depth of the concepts and the depth of their most specific common ancestor

$$sim(u, v) = depth(MSCA[u, v]) / (depth(u) + depth(v))$$
- 'edge_resnik': Computes the similarity of two concepts based on the shortest path between the concepts and the maximum depth of the taxonomy

$$(2 * max_depth - min_sp(u, v)) / (2 * max_depth)$$

max_depth is the maximum depth in the ontology
 sp(u,v) is the shortest path length between u and v
- 'edge_leachod': Computes the similarity of two concepts based on the shortest path as Rada but also considering the depth of the ontology

$$sim(u, v) = -log((sp(u, v) + 1) / 2 * max_depth)$$
- 'edge_slimani': Computes the similarity of two concepts based on the depth of the most specific common ancestor and the max depth of the concepts

$$sim(u, v) = 2 * depth(MCA) / ((depth(u) + depth(v) + 1) * pf)$$

depth(MCA) is the maximum depth of the most common ancestor of the concepts
 pf is a penalization factor used when concepts belong to the same hierarchy

The following measures require the specification of an additional measure to compute the information content of nodes.

- 'lin': Computes the similarity between two concepts based on the information content of the two concepts and the information content of the most informative common ancestor of the two concepts

$$sim(u, v) = (2 * IC(MICA)) / (IC(u) + IC(v))$$
 IC(MICA) is the information content of the most informative common ancestor of u and v. MICA is the concept in the ancestors of both u and v that maximizes the Information Content measure.
- 'resnik': Computes the similarity between two concepts based on the information content of the most informative common ancestors of the compared concepts

$$sim(u, v) = IC(MICA)$$
- 'schlicker': Computes the similarity between two concepts based on the information content of the most informative common ancestor of the compared concepts and its probability of occurrence

$$sim(u, v) = (2 * IC(MICA)) / (IC(u) + IC(v)) * (1 - Prob_{MICA})$$
 Prob_MICA is the probability of occurrence of the most informative common ancestor of the compared concepts
- 'jaccard': Computes the similarity between two concepts based on the information content of the most informative common ancestor.

$$sim(u, v) = IC(MICA) / (IC(u) + IC(v) - IC(MICA))$$
 if the sum of the IC of the concepts is different from the IC of the MICA else $sim(u, v) = 0$.
- 'sim': This measure is based on lin similarity

$$sim(u, v) = lin(u, v) - (1 - (1 / (1 + IC(MICA))))$$
- 'jc_norm': Computes the similarity between two concepts based on the IC of the most informative ancestor of the concepts

$$sim(u, v) = 1 - (IC(u) + IC(v) - 2 * IC(MICA)) / 2$$

Information content based measures require the configuration parameter for estimating concept specificity. Intrinsic estimation uses the topological properties of the taxonomic backbone of the semantic graph. There are different options:

- 'zhou': Intrinsic estimation of the specificity of the concepts based on their depth in the ontology.

$$IC(c) = k(1 - \log(D(c)) / \log(|C|)) + (1 - k)(\log(\max(\text{depth}(x))) / \log(\text{depth}_{max}))$$
 k is a factor to adjust the weight of the two items of the equation
 D(c) is the number of hyponyms of concept c
 |C| is the number of concepts in the ontology
 depth(c) is the maximum depth of concept c
 depth_max is the maximum depth in the ontology
- 'resnik_1995': Intrinsic estimation of the specificity of concepts based on the number of ancestors of the concept.

$$IC(c) = |A(c)|$$
- 'seco': Intrinsic estimation of the specificity of the concepts based on the number of concepts they subsume.

$$IC(c) = 1 - (\log(D(c)) / \log(|C|))$$
 D(c) is the number of hyponyms of concept c
 |C| is the number of concepts in the ontology

- 'sanchez': Intrinsic estimation of the specificity of the concepts based on the number of leaves and the number of subsumers of the concepts
 $IC(c) = -\log(x/nb_{leaves} + 1)$ with $x = |leaves(c)|/|A(c)|$
 nb_leaves is the represents the number of leaves corresponding to the root node of the hierarchy
 $leaves(c)$ is the number of leaves corresponding to the concept c
 $|A(c)|$ is the number of concepts that subsume c
- 'anc_norm': Intrinsic estimation of the specificity of concepts based on the number of ancestors of a given concept normalized on the number of concepts in the ontology.
- 'depth_min_non_linear': Intrinsic estimation of the specificity of concepts based on their minimum depth.
- 'depth_max_non_linear': Intrinsic estimation of the specificity of concepts based on their maximum depth.

Value

The pairwise measure

instance of the Similarity class with the new pairwise option.

Examples

```
sim <- new('Similarity')
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
ontology(sim) <- obo
pairwiseConfig(sim)
sim <- new('Similarity')
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
ontology(sim) <- obo
pairwiseConfig(sim) <- 'edge_resnik'
#The following configuration uses an information content based measure
pairwiseConfig(sim) <- c('resnik', 'seco')
```

pairwiseConfigRef	pairwiseConfigRef
-------------------	-------------------

Description

This method retrieves the reference to the Java configuration used to compute semantic similarities. configures the pairwise java object to compute semantic similarity between two concepts of a given ontology, by passing as input the java reference to one of the allowed pairwise semantic similarity measures. For a complete list check the details section of the function pairwiseConfig.

Usage

```
pairwiseConfigRef(object)

pairwiseConfigRef(object) <- value

## S4 method for signature 'Similarity'
pairwiseConfigRef(object)
```

```
## S4 replacement method for signature 'Similarity'
pairwiseConfigRef(object) <- value
```

Arguments

object	instance of class Similarity-class
value	See details

Value

The reference to the pairwise configuration used to compute semantic similarity

Examples

```
sim <- new('Similarity')
pairwiseConfigRef(sim)
sim <- new('Similarity')
pairwiseConfigRef(sim) <- c('lin')
```

paramValueIndex	<i>Method paramValueIndex</i>
-----------------	-------------------------------

Description

Method to get and set the parameter combination index corresponding to a given parameter combination. The value of the paramValueIndex lays in the range [0:575]

Usage

```
paramValueIndex(x)

paramValueIndex(x) <- value

## S4 method for signature 'COptions'
paramValueIndex(x)

## S4 replacement method for signature 'COptions'
paramValueIndex(x) <- value
```

Arguments

x	instance of class COptions
value	Index corresponding to a given parameter combination

Value

The paramValueIndex corresponding to the current options when used as get, the new COptions object with updated parameters when used to set.

Examples

```

opts <- COptions()
paramValueIndex(opts)

opts <- COptions()
paramValueIndex(opts) <- 2

```

samplesim	samplesim
-----------	-----------

Description

This method computes the semantic similarity between two named samples annotated with a group of ontology terms belonging to the same ontology

Usage

```

samplesim(object, sample1, sample2, annotated_df)

## S4 method for signature 'Similarity,character,character,data.frame'
samplesim(object,
  sample1, sample2, annotated_df)

```

Arguments

object	instance of class Similarity-class
sample1	A sample ID with its annotations available in a data frame
sample2	A sample ID with its annotations available in a data frame
annotated_df	data frame with annotations obtained using <code>entityFinder</code> . The data frame should have at least a column named 'sample_id' with the sample identifier and a column named 'term_url' with the URL of the ontology terms annotating the sample. The ontology terms must belong to the ontology loaded in the <code>Similarity</code> class.

Value

The semantic similarity between the samples `sample1` and `sample2`

Examples

```

sim <- new('Similarity')

pairwiseConfig(sim) <- listSimilarities()$pairwiseMeasures[9]
groupConfig(sim) <- listSimilarities()$groupwiseMeasures[3]
ef <- new('EntityFinder')
opts <- COptions()
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
ontology(sim) <- obo
sample_dict <- CMdictionary(inputFileOrDb=obo, outputDir=getwd(), synonymType='ALL')
sra_chip_seq <- readRDS(system.file('extdata', 'vignette_data', 'GEO_human_chip.rds',
  package='Onassis'))
chipseq_dict_annot <- EntityFinder(sra_chip_seq[1:20],c('sample_accession', 'title',

```

```
'experiment_attribute', 'sample_attribute', 'description']], dictionary=sample_dict,
options=opts)
s <- samplesim(sim, as.character(as.vector(chipseq_dict_annot$sample_id[1])),
as.character(as.vector(chipseq_dict_annot$sample_id[7])) , chipseq_dict_annot)
```

scores	scores<-
--------	----------

Description

This method gets and sets the scores slot of a class [Onassis-class](#)

Usage

```
scores(object)
```

```
scores(object) <- value
```

```
## S4 replacement method for signature 'Onassis'
scores(object) <- value
```

```
## S4 method for signature 'Onassis'
scores(object)
```

Arguments

object instance of class [Onassis-class](#)

value a matrix of scores

Value

the matrix of scores in case of get and the new [Onassis-class](#) object in case of set

Examples

```
onassis <- Onassis()
scores(onassis) <- matrix()
```

```
o <- Onassis()
scores(o)
```

SearchStrategy	<i>Method SearchStrategy</i>
----------------	------------------------------

Description

Method to get and set the SearchStrategy parameter

Usage

```
SearchStrategy(x)

SearchStrategy(x) <- value

## S4 method for signature 'COptions'
SearchStrategy(x)

## S4 replacement method for signature 'COptions'
SearchStrategy(x) <- value
```

Arguments

x	instance of class COptions
value	The matching strategy for finding concepts in the input text <ul style="list-style-type: none">• CONTIGUOUS_MATCH Longest match of contiguous tokens within enclosing span• SKIP_ANY_MATCH Longest match of not-necessarily contiguous tokens• SKIP_ANY_MATCH_ALLOW_OVERLAP Longest match of not-necessarily contiguous tokens, overlapping matches are allowed

Value

The SearchStrategy corresponding to the current options when used as get, the new COptions object with updated parameters when used to set.

Examples

```
opts <- COptions()
SearchStrategy(opts)

opts <- COptions()
SearchStrategy(opts) <- 'SKIP_ANY_MATCH_ALLOW_OVERLAP'
```

```
semanticdifference    semanticdifference
```

Description

semanticdifference function allows to test, within a given tissue class, the difference between i) the healthy samples and one or more disease states and ii) if there is a significant difference between the different disease states for each unit represented in each row of a scorematrix containing as rows genomic units and as columns GEO sample ids.

Usage

```
semanticdifference(score_matrix, list_of_annotations, fun_name, test_type)
```

Arguments

score_matrix	A matrix where rows represent units (GRanges or genes) and columns #' represent GSMs.
list_of_annotations	A list of lists where the first level elements represent a tissue/cell line semantic set, the second level elements represent disease semantic sets and each element of the list is a named score matrix with column names corresponding to sample (GSM) identifiers
fun_name	The name of a testing function to measure the differences between semantic classes. For the test_type parameter = 'pair' the semanticdifference function applies the test function to the tissue semantic states including 'Healthy' and one or more diseases. In this case the function (e.g. wilcoxon.test) should take as input a couple of vectors and return as output the value of the of the statistic '\$statistic' field and a p-value '\$p.value' field. For the test_type = 'multiple' the function tests if there are differences between the semantic disease classes within the tissue class. The function (e.g. kruskal.test) takes as input a list of vectors (one for each disease semantic set) and returns the value of the statistic in the '\$statistic' field and the corresponding p-value.
test_type	This value can be 'pair' or 'multiple'. In case it is set to pair the function to provide in fun_name requires the tissue semantic class to include at least two disease semantic classes of which one has to be 'Healthy' to compare diseases against healthy. For example the 'wilcox.test'. When set to multiple the fun_name provided should take as input a list of input vectors corresponding to the different disease conditions. A minimum of three is required. For example the 'kruskal.test' can be used.

Value

The semanticdifference function returns a list where each element corresponds to a tissue/cell line semantic state. The content of the list for 'multiple' test types a matrix of the test results and corresponding p.values.

Examples

```
granges <- readRDS(system.file('extdata', 'sample_granges.rds', package='Onassis'))
ref_granges <- granges[[1]]
for(i in 2:length(granges)) {
  ref_granges <- GenomicRanges::union(ref_granges, granges[[i]])
}
score_mat <- create_score_matrix(ref_granges, granges)
gsm_list <- names(granges)
list_of_annotations <- readRDS(system.file('extdata', 'list_of_annotations.rds', package='Onassis'))
fun_name = 'wilcox.test'
fun_type = 'pair'
sem_dif <- semanticdifference(score_mat, list_of_annotations, fun_name, fun_type)
```

sim

sim

Description

This method computes the similarities of the entities annotated in a object fo class [Onassis-class](#).

Usage

```
sim(onassis = NA, ...)

## S4 method for signature 'Onassis'
sim(onassis, iconf = "sanchez", pairconf = "lin",
    groupconf = "bma")
```

Arguments

onassis	instance of class Onassis-class
...	Optional parameters
iconf	the information content measure see <code>pairwiseConfig</code> help for details
pairconf	the pairwise measure to compute semantic similarity between single concepts. See <code>pariwiseConfig</code> help for details
groupconf	the groupwise measure to compute semantic similarity between groups of concepts. See <code>groupConfig</code> help for details

Value

an instance of [Onassis-class](#) with computed similarities

Examples

```
geo_chip <- readRDS(system.file('extdata', 'vignette_data', 'GEO_human_chip.rds', package='Onassis'))
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
onassis_results <- annotate(geo_chip, 'OBO', dictionary=obo)
entities <- entities(onassis_results)
entities(onassis_results) <- entities[sample(nrow(entities), 30),]
onassis_results <- sim(onassis_results)
```

simil	simil<-
-------	---------

Description

Method to get and set the similarity slot of the class `Onassis-class`

Usage

```
simil(object)
```

```
simil(object) <- value
```

```
## S4 replacement method for signature 'Onassis'
simil(object) <- value
```

```
## S4 method for signature 'Onassis'
simil(object)
```

Arguments

`object` instance of class `Onassis-class`

`value` a matrix of similarities between the entities of an object of class `Onassis-class`

Value

the similarity matrix of an object of type `Onassis-class` in case of get, the new `Onassis-class` instance in case of set

Examples

```
onassis <- Onassis()
simil(onassis) <- matrix()
```

```
o <- Onassis()
simil(o)
```

Similarity	Similarity
------------	------------

Description

this constructr initializes the Similarity class to compute the similarity between couple of terms, couple of samples, or group of terms

Usage

```
Similarity(ontology, termlist1, termlist2, annotatedtab = NA,
  icConf = "seco", pairConf = "lin", groupConf = "ui")
```


Arguments

ontology	The ontology file to create the DAG to compute similarities
termlist1	The single concept or vector of concepts in the first set or the name of a sample if annotatedtab contains an annotated table
termlist2	The single concept or vector of concepts in the second set or the name of a sample if annotatedtab contains an annotated table
annotatedtab	The table of annotation of samples or entities
icConf	the information content configuration
pairConf	the pairwise configuration
groupConf	the groupwise configuration

Value

Measure of the similarity bewtween the concepts passed as input

Examples

```
obo <- system.file('extdata', 'sample.cs.obo', package='OnassisJavaLibs')
sample_dict <- CMdictionary(input=obo, outputDir=getwd(), synonymType='ALL')
myopts <- COptions(paramValueIndex=40)
term_list1 <- c('http://purl.obolibrary.org/obo/CL_0000000', 'http://purl.obolibrary.org/obo/CL_0000236')
term_list2 <- c('http://purl.obolibrary.org/obo/CL_0000542')
sim <- Similarity(obo, term_list1, term_list2)
```

Similarity-class	<i>Similarity class to compute similarities between concepts in ontologies and samples annotated with different concepts</i>
------------------	--

Description

Similarity is a class that wraps some methods of the Java library <http://www.semantic-measures-library.org/sml/>. Starting from OBO ontologies it is possible to build semantic graphs that allow the computation of different similarity measures between concepts belonging to the same ontology, group of concepts, samples annotated with different ontology concepts. Further details about the graph based semantic similarity measures are available at http://www.semantic-measures-library.org/sml/index.php?q=doc_graph_based_advanced

Details

The following methods can be applied to Similarity

```
ontology<-
pairwiseConfig
groupConfig
sim
groupsim
samplesim
```

To see the available similarity measures run the function `listSimilarities`

Slots

similarityInstance The Java reference to the Java Similarity class.

pairwiseConfig The list of measures used to compute the semantic similarity between two concepts in the same ontology.

pairwiseConfigRef The reference to the Java object of type CMconf corresponding to the pairwise configuration

groupConfig The groupwise configuration to compute the semantic similarity between groups of concepts.

icConfig The information content measure

groupwiseConfigRef The reference to the Java configuration object for the computation of semantic similarity between groups of concepts

ontology The ontology to compute semantic similarities

Examples

```
sim <- new('Similarity')
```

```
similarityInstance    similarityInstance
```

Description

This method retrieves the java object referencing the Similarity class

Usage

```
similarityInstance(object)

similarityInstance(object) <- value

## S4 method for signature 'Similarity'
similarityInstance(object)

## S4 replacement method for signature 'Similarity'
similarityInstance(object) <- value
```

Arguments

object	Instance of the class Similarity-class
value	the reference to a java Similarity object

Value

The java reference to an object of class Similarity

The object of class Similarity with a new instance of the java Similarity class

Examples

```
sim <- new('Similarity')
similarityInstance(sim)
sim <- new('Similarity')
similarityInstance(sim) <- .jnew('iit/comp/epigen/nlp/similarity/Similarity')
```

 Stemmer

Method Stemmer

Description

Method to get and set the Stemmer parameter

Usage

```
Stemmer(x)

Stemmer(x) <- value

## S4 method for signature 'COptions'
Stemmer(x)

## S4 replacement method for signature 'COptions'
Stemmer(x) <- value
```

Arguments

x	instance of class COptions
value	<ul style="list-style-type: none"> • BIOLEMMATIZER A stemmer specific for biomedical literature • PORTER A stemmer that removes the commoner morphological and inflexional endings from words in English • NONE No word stemming

Value

The Stemmer corresponding to the current options when used as get, the new COptions object with updated parameters when used to set.

Examples

```
opts <- COptions()
Stemmer(opts)

opts <- COptions()
Stemmer(opts) <- 'PORTER'
```

 StopWords

Method StopWords

Description

Method to get and set the StopWords parameter

Usage

```
StopWords(x)
```

```
StopWords(x) <- value
```

```
## S4 method for signature 'COptions'  
StopWords(x)
```

```
## S4 replacement method for signature 'COptions'  
StopWords(x) <- value
```

Arguments

x	instance of class COptions
value	<ul style="list-style-type: none"> • PUBMED A list of stop words obtained analyzing Pubmed papers • NONE No stop words

Value

The StopWords corresponding to the current options when used as get, the new COptions object with updated parameters when used to set.

Examples

```
opts <- COptions()  
StopWords(opts)
```

```
opts <- COptions()  
StopWords(opts) <- 'NONE'
```

 SynonymType

Method SynonymType

Description

Method to get and set the SynonymType parameter

Usage

```

SynonymType(x)

SynonymType(x) <- value

## S4 method for signature 'CMoptions'
SynonymType(x)

## S4 replacement method for signature 'CMoptions'
SynonymType(x) <- value

```

Arguments

`x` instance of class [CMoptions](#)

`value`

- EXACT_ONLY Only exact synonyms are considered
- ALL All synonym types are included

Value

The SynonymType corresponding to the current options when used as get, the new CMoptions object with updated parameters when used to set.

Examples

```

opts <- CMoptions()
SynonymType(opts)

opts <- CMoptions()
SynonymType(opts) <- 'ALL'

```

typeSystemRef	<i>Method typeSystemRef</i>
---------------	-----------------------------

Description

This method sets the type system to the ccp-nlp one to run the EntityFinder

This method sets the type system to the ccp-nlp one to run the EntityFinder

Usage

```

typeSystemRef(x)

typeSystemRef(x) <- value

## S4 method for signature 'EntityFinder'
typeSystemRef(x)

## S4 replacement method for signature 'EntityFinder'
typeSystemRef(x) <- value

```

Arguments

x instance of the class EntityFinder
value the java type system to detect concepts from ontologies.

Value

the reference to the Java type system currently set
The updated EntityFinder S4 object

Examples

```
ef <- new('EntityFinder')
typeSystemRef(ef)
ef <- new('EntityFinder')
type_system_array_list <- .jnew('java/util/ArrayList')
ccp_nlp_type_system <- .jfield('edu/ucdenver/ccp/nlp/uima/util/TypeSystemUtil',
  name = 'CCP_TYPE_SYSTEM')
sentence_detector_type_system_str <- 'org.cleartk.token.type.Sentence'
conceptmapper_type_system <-
'edu.ucdenver.ccp.nlp.wrapper.conceptmapper.TypeSystem'
dictTerm <- 'analysis_engine.primitive.DictTerm'
tokenizer <- 'org.apache.uima.conceptMapper.support.tokenizer.TokenAnnotation'
vector_of_ts <- c(ccp_nlp_type_system, sentence_detector_type_system_str,
  conceptmapper_type_system, dictTerm, tokenizer)
type_system_description <-
J('org/uimafit/factory/TypeSystemDescriptionFactory')$createTypeSystemDescription(vector_of_ts)
typeSystemRef(ef) <- type_system_description
```

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