

Package ‘epos’

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Type Package

Title Epilepsy Ontologies' Similarities

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Description Analysis and visualization of similarities between epilepsy ontologies based on text mining results by comparing ranked lists of co-occurring drug terms in the BioASQ corpus. The ranked result lists of neurological drug terms co-occurring with terms from the epilepsy ontologies EpSO, ESSO, EPILONT, EPISEM and FEN-ICS undergo further analysis. The source data to create the ranked lists of drug names is produced using the text mining workflows described in Mueller, Bernd and Hagelstein, Alexandra (2016) <[doi:10.4126/FRL01-006408558](https://doi.org/10.4126/FRL01-006408558)>, Mueller, Bernd et al. (2017) <[doi:10.1007/978-3-319-58694-6_22](https://doi.org/10.1007/978-3-319-58694-6_22)>, Mueller, Bernd and Rebholz-Schuhmann, Dietrich (2020) <[doi:10.1007/978-3-030-43887-6_52](https://doi.org/10.1007/978-3-030-43887-6_52)>, and Mueller, Bernd et al. (2022) <[doi:10.1186/s13326-021-00258-w](https://doi.org/10.1186/s13326-021-00258-w)>.

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License LGPL (>= 3)

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URL <https://github.com/bernd-mueller/epos>

BugReports <https://github.com/bernd-mueller/epos/issues>

Imports hash, ggplot2, testthat, gridExtra, TopKLists, stringr, xtable, mongolite, stats, VennDiagram, cowplot

Suggests knitr, rmarkdown

NeedsCompilation no

Repository CRAN

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calcCosine

Calculate the cosine similarity metric for two lists a and b

Description

Calculate the cosine similarity metric for two lists a and b

Usage

```
calcCosine(a, b)
```

Arguments

- | | |
|---|---|
| a | list with elements that should be of same type as in list b |
| b | list with elements |

Value

co list with length of set b containing the cosine similarity coefficient at each position

Examples

```
calcCosine(c(1,2), c(2,3))
```

calcDice

Calculate the dice similarity metric for two lists a and b

Description

Calculate the dice similarity metric for two lists a and b

Usage

```
calcDice(a, b)
```

Arguments

- | | |
|---|---|
| a | list with elements that should be of same type as in list b |
| b | list with elements |

Value

di list with length of set b containing the dice similarity coefficient at each list element

Examples

```
calcDice(c(1,2), c(2,3))
```

calcDSEA*Calculate dsea scores of one list in comparison to reference list***Description**

Calculate dsea scores of one list in comparison to reference list

Usage

```
calcDSEA(alist, N)
```

Arguments

| | |
|-------|---|
| alist | list of drug names to be used for calculating dsea |
| N | numeric value with maximum length of lists for dsea calculation |

Value

list with dsea scores

Examples

```
calcDSEA(c("Valproic acid", "Lamotrigine", "Ketamin"), 3)
```

calcEnrichment*Calculate enrichment of one list in comparison to reference list***Description**

Calculate enrichment of one list in comparison to reference list

Usage

```
calcEnrichment(alist)
```

Arguments

| | |
|-------|---------------------|
| alist | the list to compare |
|-------|---------------------|

Value

list with calculated enrichment used for plotting

Examples

```
a <- calcEnrichment(c("Clobazam", "Oxcarbazepine"))
```

calcJaccard*Calculate the jaccard coefficient for two lists a and b*

Description

Calculate the jaccard coefficient for two lists a and b

Usage

```
calcJaccard(a, b)
```

Arguments

| | |
|---|---|
| a | list with elements that should be of same type as in list b |
| b | list with elements |

Value

a list with length of set b containing the jaccard similarity coefficient for each list element

Examples

```
calcJaccard(c(1,2), c(2,3))
```

cosine*Calculate cosine similarity metric*

Description

Calculate cosine similarity metric

Usage

```
cosine(ainterb, lengtha, lengthb)
```

Arguments

| | |
|---------|--|
| ainterb | integer value with number of intersecting elements between set a and b |
| lengtha | integer value with the number of items in set a |
| lengthb | integer value with the number of items in set b |

Value

cosine double vlaue with the cosine similarity coefficient

Examples

```
cosine(1,3,4)
```

| | |
|------------------------------|---|
| <code>createBaseTable</code> | <i>Main function to call everything and produce the results</i> |
|------------------------------|---|

Description

Main function to call everything and produce the results

Usage

```
createBaseTable(coocepso, coocesso, coocepi, coocepisem, coocfenics)
```

Arguments

| | |
|------------|---|
| coocepso | list of drug names sorted by frequency co-occuring with EpSO |
| coocesso | list of drug names sorted by frequency co-occuring with ESSO |
| coocepi | list of drug names sorted by frequency co-occuring with EPILONT |
| coocepisem | list of drug names sorted by frequency co-occuring with EPISEM |
| coocfenics | list of drug names sorted by frequency co-occuring with FENICS |

Value

result table containin the aggregated list of drug terms and their associations

Examples

```
utils::data(rawDrugNamesCo0cEpSO, package="epos")
utils::data(rawDrugNamesCo0cESSO, package="epos")
utils::data(rawDrugNamesCo0cEPILONT, package="epos")
utils::data(rawDrugNamesCo0cEPISEM, package="epos")
utils::data(rawDrugNamesCo0cFENICS, package="epos")
createBaseTable(coocepso = rawDrugNamesCo0cEpSO[1:150],
               coocesso=rawDrugNamesCo0cESSO[1:150],
               coocepi=rawDrugNamesCo0cEPILONT[1:150],
               coocepisem=rawDrugNamesCo0cEPISEM[1:150],
               coocfenics=rawDrugNamesCo0cFENICS[1:150])
```

| | |
|-------------------------------------|--|
| <code>createDashVectorForATC</code> | <i>Creates a vector with an X at each position where a drug from the druglist matches the ATC class list slate</i> |
|-------------------------------------|--|

Description

Creates a vector with an X at each position where a drug from the druglist matches the ATC class list slate

Usage

```
createDashVectorForATC(druglist, atchashda, atchashsec, slatc)
```

Arguments

| | |
|------------|--|
| druglist | list of drug names |
| atchashda | hash retrieved from readAtcMapIntoHashMapDrugNamesAtcCodes |
| atchashsec | hash retrieved from readSecondLevelATC |
| slatc | list of ATC classes |

Value

list with crosses if the drug in druglist matches at the position of the ATC class in slatc

Examples

```
## Not run:  
createDashVectorForATC(druglist, atchashda, atchashsec, slatc)  
  
## End(Not run)
```

createJaccardPlotDBMeSH

Creates the plot for all jaccard coefficients amongst the three epilepsy ontologies

Description

Creates the plot for all jaccard coefficients amongst the three epilepsy ontologies

Usage

```
createJaccardPlotDBMeSH(jmeshepso, jmehessso, jmeshepi)
```

Arguments

| | |
|-----------|---|
| jmeshepso | list containing jaccard coefficients between mesh and epso for increasing k |
| jmehessso | list containing jaccard coefficients between mesh and esso for increasing k |
| jmeshepi | list containing jaccard coefficients between mesh and epi for increasing k |

Value

jaccardepilepsyplot the ggplot object

Examples

```
## Not run:
jaccardepilepsyplot <- createJaccardPlotAll(jaccardepso, jaccardesso)

## End(Not run)
```

createJaccardPlotMeSHFive

Creates the plot for all jaccard coefficients amongst the three epilepsy ontologies

Description

Creates the plot for all jaccard coefficients amongst the three epilepsy ontologies

Usage

```
createJaccardPlotMeSHFive(
  jmeshepso,
  jmeshesso,
  jmeshepi,
  jmeshepilepsyand,
  jmeshepilepsyor
)
```

Arguments

| | |
|------------------|--|
| jmeshepso | list of jaccard coefficients between mesh and epso for increasing k |
| jmeshesso | list of jaccard coefficients between mesh and esso for increasing k |
| jmeshepi | list of jaccard coefficients between mesh and epi for increasing k |
| jmeshepilepsyand | list of jaccard coefficients between mesh and the intersection of epso, esso, and epi for increasing k |
| jmeshepilepsyor | list of jaccard coefficients between mesh and the union of epso, esso, and epi for increasing k |

Value

jaccardepilepsyplot the ggplot object

Examples

```
## Not run:
jaccardepilepsyplot <- createJaccardPlotAll(jaccardepso, jaccardesso)

## End(Not run)
```

| | |
|-------------------------------|--|
| <code>createNeuroTable</code> | <i>Create the final resulting data frame</i> |
|-------------------------------|--|

Description

Create the final resulting data frame

Usage

```
createNeuroTable(atchashda, atchashsec, dneuromaxk)
```

Arguments

| | |
|------------|--|
| atchashda | hashmap retrieved from <code>readAtcMapIntoHashMapDrugNamesAtcCodes</code> |
| atchashsec | hashmap retrieved from <code>readSecondLevelATC</code> |
| dneuromaxk | data frame containing columns for each intersection, ATC class, and reference list |

Value

data frame containing drug names with additional columns listing association to ATC classes

Examples

```
utils::data(rawDrugNamesCo0cEpSO, package="epos")
utils::data(rawDrugNamesCo0cESSO, package="epos")
utils::data(rawDrugNamesCo0cEPILONT, package="epos")
utils::data(rawDrugNamesCo0cEPISEM, package="epos")
utils::data(rawDrugNamesCo0cFENICS, package="epos")
atchashda <-
readAtcMapIntoHashMapDrugNamesAtcCodes(
  system.file("extdata", "db-atc.map", package = "epos"), "\t")
atchashaa <-
readAtcMapIntoHashMapAtcCodesAtcNames(
  system.file("extdata", "db-atc.map", package = "epos"), "\t")
atchashsec <-
readSecondLevelATC(
  system.file("extdata", "atc-secondlevel.map", package = "epos"), "\t")
epso <- rawDrugNamesCo0cEpSO
neuroepso <- filterNeuroDrugs(epso, atchashda)
esso <- rawDrugNamesCo0cESSO
neuroesso  <- filterNeuroDrugs(esso, atchashda)
epi <- rawDrugNamesCo0cEPILONT
neuroepi   <- filterNeuroDrugs(epi, atchashda)
episem <- rawDrugNamesCo0cEPISEM
neuroepisem <- filterNeuroDrugs(episem, atchashda)
fenics <- rawDrugNamesCo0cFENICS
neurofenics <- filterNeuroDrugs(fenics, atchashda)
mx <- max(
```

```

c(length(neuroepso), length(neuroesso), length(neuroepi),
  length(neuroepisem), length(neurofenics)))
dneuro <-
  data.frame(EpSO = c(neuroepso, rep(1, (mx-length(neuroepso)))),
             ESSO = c(neuroesso, rep(1, (mx-length(neuroesso)))),
             EPILONT = c(neuroepi, rep(1, (mx-length(neuroepi)))),
             EPISEM = c(neuroepisem, rep(1, (mx-length(neuroepisem)))),
             FENICS = c(neurofenics, rep(1, (mx-length(neurofenics)))))

dneuromaxk <- TopKLists::calculate.maxK(dneuro, L=5, d=5, v=10)
neurotable <- createNeuroTable(atchashda, atchashsec, dneuromaxk)

```

createTanimotoBaseline

Creates the plot for all jaccard coefficients amongst the three epilepsy ontologies

Description

Creates the plot for all jaccard coefficients amongst the three epilepsy ontologies

Usage

```
createTanimotoBaseline(neuroepso, neuroesso, neuroepi, dneuromaxk)
```

Arguments

| | |
|------------|---|
| neuroepso | list of neuro drug names co-occurring with epso |
| neuroesso | list of neuro drug names co-occurring with esso |
| neuroepi | list of neuro drug names co-occurring with epi |
| dneuromaxk | object returned from TopKLists::calculate.maxk |

Value

jaccardepilepsyplot the ggplot object

Examples

```

utils::data(rawDrugNamesCo0cEpSO, package="epos")
utils::data(rawDrugNamesCo0cESSO, package="epos")
utils::data(rawDrugNamesCo0cEPILONT, package="epos")
utils::data(rawDrugNamesCo0cEPISEM, package="epos")
atchashda <-
  readAtcMapIntoHashMapDrugNamesAtcCodes(
    system.file("extdata", "db-atc.map", package = "epos"), "\t")
atchashaa <-
  readAtcMapIntoHashMapAtcCodesAtcNames(
    system.file("extdata", "db-atc.map", package = "epos"), "\t")
atchashsec <

```

```

readSecondLevelATC(
  system.file("extdata", "atc-secondlevel.map", package = "epos"), "\t")
tepso <- rawDrugNamesCoOcEpSO
tesso <- rawDrugNamesCoOcESSO
tepi <- rawDrugNamesCoOcEPILONT
tepsem <- rawDrugNamesCoOcEPISEM
tfenics <- rawDrugNamesCoOcFENICS

neuroepso <- filterNeuroDrugs(tepso, atchashda)
neuroesso <- filterNeuroDrugs(tesso, atchashda)
neuroepi <- filterNeuroDrugs(tepi, atchashda)
neuroepisem <- filterNeuroDrugs(tepsem, atchashda)
neurofenics <- filterNeuroDrugs(tfenics, atchashda)

dneuro <-
  data.frame(EpSO = neuroepso[1:210],
             ESSO = neuroesso[1:210],
             EPILONT = neuroepi[1:210],
             EPISEM = neuroepisem[1:210],
             FENICS = neurofenics[1:210])
dneuromaxk <- TopKLists::calculate.maxK(dneuro, 5, 5, 5)
tanimotobaseline <- createTanimotoBaseline(neuroepso, neuroesso, neuroepi, dneuromaxk)

```

dice

*Calculate dice similarity metric***Description**

Calculate dice similarity metric

Usage

```
dice(ainterb, lengtha, lengthb)
```

Arguments

| | |
|---------|--|
| ainterb | integer value with number of intersecting elements between set a and b |
| lengtha | integer value with the number of items in set a |
| lengthb | integer value with the number of items in set b |

Value

dice double vlaue with the dice similarity coefficient

Examples

```
dice(1, 3, 4)
```

| | |
|------------|---------------------------------------|
| doFullPlot | <i>Does the full plot on one page</i> |
|------------|---------------------------------------|

Description

Does the full plot on one page

Usage

```
doFullPlot(
  cosinemeshplot,
  cosinedrugbankplot,
  cosinepilepsyplot,
  dicemeshplot,
  dicedrugbankplot,
  dicepilepsyplot,
  jaccardmeshplot,
  jaccarddrugbankplot,
  jaccardepilepsyplot
)
```

Arguments

| | |
|---------------------|---|
| cosinemeshplot | plot with cosine coefficients against MeSH |
| cosinedrugbankplot | plot with cosine coefficients against DrugBank |
| cosinepilepsyplot | plot with cosine coefficients of Epilepsy Ontologies |
| dicemeshplot | plot with dice coefficients against MeSH |
| dicedrugbankplot | plot with dice coefficients against DrugBank |
| dicepilepsyplot | plot with dice coefficients of Epilepsy Ontologies |
| jaccardmeshplot | plot with jaccard coefficients against MeSH |
| jaccarddrugbankplot | plot with jaccard coefficients against DrugBank |
| jaccardepilepsyplot | plot with jaccard coefficients of Epilepsy Ontologies |

Value

full

Examples

```
## Not run:  
full <- doFullPlot (cosinemeshplot,  
                      cosinedrugbankplot,  
                      cosinepilepsyplot,  
                      dicemeshplot,  
                      dicedrugbankplot,  
                      dicepilepsyplot,  
                      jaccardmeshplot,  
                      jaccarddrugbankplot,  
                      jaccardepilepsyplot)  
  
## End(Not run)
```

drawVenn4

*Create quad Venn Diagramm for overlapping concepts between EpSO,
ESSO, EPILONT and EPISEM*

Description

Create quad Venn Diagramm for overlapping concepts between EpSO, ESSO, EPILONT and EPISEM

Usage

```
drawVenn4()
```

Value

plot object

Examples

```
## Not run:  
ggplot2::ggsave("venn4.png", plot = drawVenn4(), width=240, height=160,  
                 units = "mm", dpi = 300)  
  
## End(Not run)
```

drawVenn4Doc

Create quintuple Venn Diagramm for shared documents with co-occurrences of drug names between EpSO, ESSO, EPILONT and EPISEM

Description

Create quintuple Venn Diagramm for shared documents with co-occurrences of drug names between EpSO, ESSO, EPILONT and EPISEM

Usage

```
drawVenn4Doc()
```

Value

plot object

Examples

```
## Not run:  
ggplot2::ggsave("venn4doc.png", plot = drawVenn4Doc(), width=240, height=160,  
units = "mm", dpi = 300)  
  
## End(Not run)
```

drawVenn4DrugDoc

Create quad Venn Diagramm for shared documents with co-occurrences of drug names between EpSO, ESSO, EPILONT and EPISEM

Description

Create quad Venn Diagramm for shared documents with co-occurrences of drug names between EpSO, ESSO, EPILONT and EPISEM

Usage

```
drawVenn4DrugDoc()
```

Value

plot object

Examples

```
## Not run:
ggplot2::ggsave("venn4drugdoc.png", plot = drawVenn4DrugDoc(), width=240,
height=160, units = "mm", dpi = 300)

## End(Not run)
```

drawVenn4Syn

*Create quad Venn Diagramm for shared synonyms between EpSO,
ESSO, EPILONT and EPISEM*

Description

Create quad Venn Diagramm for shared synonyms between EpSO, ESSO, EPILONT and EPISEM

Usage

drawVenn4Syn()

Value

plot object

Examples

```
## Not run:
ggplot2::ggsave("venn4syn.png", plot = drawVenn4Syn(), width=240,
height=160, units = "mm", dpi = 300)

## End(Not run)
```

drawVenn5

*Create quintuple Venn Diagramm for overlapping concepts between
EpSO, ESSO, EPILONT, EPISEM and FENICS*

Description

Create quintuple Venn Diagramm for overlapping concepts between EpSO, ESSO, EPILONT, EPISEM and FENICS

Usage

drawVenn5()

Value

plot object

Examples

```
## Not run:
ggplot2::ggsave("venn5.png", plot = drawVenn5(), width=240, height=160,
  units = "mm", dpi = 300)

## End(Not run)
```

drawVenn5Doc

Create quintuple Venn Diagramm for shared documents between EpSO, ESSO, EPILONT, EPISEM and FENICS

Description

Create quintuple Venn Diagramm for shared documents between EpSO, ESSO, EPILONT, EPISEM and FENICS

Usage

```
drawVenn5Doc()
```

Value

plot object

Examples

```
## Not run:
ggplot2::ggsave("venn5doc.png", plot = drawVenn5Doc(), width=240, height=160,
  units = "mm", dpi = 300)

## End(Not run)
```

drawVenn5DrugDoc

Create quintuple Venn Diagramm for shared documents with co-occurrences of drug names between EpSO, ESSO, EPILONT, EPISEM and FENICS

Description

Create quintuple Venn Diagramm for shared documents with co-occurrences of drug names between EpSO, ESSO, EPILONT, EPISEM and FENICS

Usage

```
drawVenn5DrugDoc()
```

Value

plot object

Examples

```
## Not run:  
ggplot2::ggsave("venn5drugdoc.png", plot = drawVenn5DrugDoc(), width=240,  
height=160, units = "mm", dpi = 300)  
  
## End(Not run)
```

drawVenn5Syn

*Create quintuple Venn Diagramm for shared synonyms between EpSO,
ESSO, EPILONT, EPISEM and FENICS*

Description

Create quintuple Venn Diagramm for shared synonyms between EpSO, ESSO, EPILONT, EPISEM and FENICS

Usage

drawVenn5Syn()

Value

plot object

Examples

```
## Not run:  
ggplot2::ggsave("venn5syn.png", plot = drawVenn5Syn(), width=240,  
height=160, units = "mm", dpi = 300)  
  
## End(Not run)
```

drawVennGrid

Create plot_grid from multiple plots

Description

Create plot_grid from multiple plots

Usage

drawVennGrid()

Value

plot object

Examples

```
## Not run:
cowplot::plot_grid(drawVenn4 (), drawVenn4Syn(), drawVenn5Doc (),
  drawVenn5DrugDoc ())
ggplot2::ggsave("vennAB.png", plot = cowplot::plot_grid(drawVenn4 (),
  drawVenn4Syn(), labels = c('A', 'B'), ncol = 1), width=240, height=320,
  units = "mm", dpi = 300)
ggplot2::ggsave("vennAB.png", plot = cowplot::plot_grid(drawVenn4 (),
  drawVenn4Syn(), labels = c('Concepts:', 'Synonyms:'), ncol = 1), width=240,
  height=320, units = "mm", dpi = 300)
ggplot2::ggsave("vennCD.png", plot = cowplot::plot_grid(drawVenn5Doc (),
  drawVenn5DrugDoc(), labels = c('Documents with B-Terms:',
  'Documents with B- and C-Terms:'), ncol = 1), width=240, height=320,
  units = "mm", dpi = 300)
ggplot2::ggsave("vennCD.png", plot = cowplot::plot_grid(drawVenn5Doc (),
  drawVenn5DrugDoc(), labels = c('Documents with B-Terms:',
  'Documents with B- and C-Terms:'), ncol = 1), width=240, height=320, units = "mm",
  dpi = 300)
ggplot2::ggsave("vennCD.png", plot = cowplot::plot_grid(drawVenn4Doc (),
  drawVenn4DrugDoc(), labels = c('Documents with B-Terms:',
  'Documents with B- and C-Terms:'), ncol = 1), width=240, height=320,
  units = "mm", dpi = 300)
ggplot2::ggsave("vennCD.png", plot = cowplot::plot_grid(drawVenn4Doc (),
  drawVenn4DrugDoc(), labels = c('Documents\nwith B-Terms:',
  'Documents\nwith B- and C-Terms:'), ncol = 1), width=240, height=320,
  units = "mm", dpi = 300)
ggplot2::ggsave("vennAB.png", plot = cowplot::plot_grid(drawVenn4 (),
  drawVenn4Syn(), labels = c('i) Concepts:', 'ii) Synonyms:'), ncol = 1),
  width=240, height=320, units = "mm", dpi = 300)
ggplot2::ggsave("vennCD.png", plot = cowplot::plot_grid(NULL,
  drawVenn4Doc (), drawVenn4DrugDoc(),
  labels = c('iii) Documents with B-Terms:',
  'iv) Documents with B- and C-Terms:'), ncol = 1,
  label_x = c(-0.105, -0.14), label_fontfamily = "Arial Nova Light",
  label_fontface = "bold"), width=240, height=320, units = "mm", dpi = 300)

## End(Not run)
```

filterApprovedDrugs *Filter a given list of drug names for having an ATC code, if not they are dropped*

Description

Filter a given list of drug names for having an ATC code, if not they are dropped

Usage

```
filterApprovedDrugs(druglist, atchashda)
```

Arguments

| | |
|-----------|--|
| druglist | a list of drug names |
| atchashda | a hash containing the drug names as keys |

Value

approveddrugs a hash filtered for having an ATC code

Examples

```
utils::data(rawDrugNamesCo0cEpS0, package="epos")
atchashda <-
  readAtcMapIntoHashMapDrugNamesAtcCodes(
    system.file("extdata", "db-atc.map", package = "epos"), "\t")
tepso <- genDictListFromRawFreq(rawDrugNamesCo0cEpS0)
filterApprovedDrugs(tepso, atchashda)
```

filterNeuroDrugs

Filter a given list of drug names for having an ATC code starting with N indicating to be a drug for the Nervous System

Description

Filter a given list of drug names for having an ATC code starting with N indicating to be a drug for the Nervous System

Usage

```
filterNeuroDrugs(druglist, atchashda)
```

Arguments

| | |
|-----------|--|
| druglist | a list of drug names |
| atchashda | a hash containing the drug names as keys |

Value

neurodrugs a hash filtered for having an ATC code starting with N

Examples

```
utils::data(rawDrugNamesCo0cEpS0, package="epos")
atcHashda <-
  readAtcMapIntoHashMapDrugNamesAtcCodes(
    system.file("extdata", "db-atc.map", package = "epos"), "\t")
tepso <- genDictListFromRawFreq(rawDrugNamesCo0cEpS0)
nepso <- filterNeuroDrugs(tepso, atcHashda)
```

genDictListFromRawFreq

Clears object that was loaded from harddrive into a list of terms sorted by frequency

Description

Clears object that was loaded from harddrive into a list of terms sorted by frequency
 Clears object that was loaded from harddrive into a list of terms sorted by frequency

Usage

```
genDictListFromRawFreq(topfreqdictraw)
genDictListFromRawFreq(topfreqdictraw)
```

Arguments

`topfreqdictraw` list with terms from a dictionary sorted by frequency

Value

a sorted list of terms
 a sorted list of terms

Examples

```
## Not run:
genDictListFromRawFreq(epi)

## End(Not run)
utils::data(rawDrugNamesCo0cEpS0, package="epos")
genDictListFromRawFreq(rawDrugNamesCo0cEpS0)
```

getRefAll*Retrieve the list of drugs from the union of all reference lists*

Description

Retrieve the list of drugs from the union of all reference lists

Usage

```
getRefAll()
```

Value

list of drugs from all reference lists

Examples

```
d <- getRefAll()
```

getTermMatrix*Receives a sorted hashmap with found entities from a dictionary*

Description

Receives a sorted hashmap with found entities from a dictionary

Usage

```
getTermMatrix(dictionary, database, collection)
```

Arguments

| | |
|------------|---|
| dictionary | Character vector that is the name of a dicitonary having pre-calculated stats. This can be MeSH, DrugBank, Agrovoc, EpSO, ESSO, or EPILONT |
| database | the name of the MongoDB database to be used |
| collection | the name of the MongoDB collection to be used |

Value

a sorted hashmap containing all found entities from the respective dictionaries with frequencies

Examples

```
## Not run:  
mesh <- getTermMatrix("MeSH")  
  
## End(Not run)
```

jaccard*Calculate jaccard similarity metric for two sets a and b***Description**

Calculate jaccard similarity metric for two sets a and b

Usage

```
jaccard(ainterb, aunionb, lengtha, lengthb)
```

Arguments

| | |
|---------|--|
| ainterb | integer value with number of intersecting elements between set a and b |
| aunionb | integer value with number of union elements between set a and b |
| lengtha | length of set a |
| lengthb | length of set b |

Value

jac double value with the jaccard similarity coefficient

Examples

```
jaccard(1,3, 2, 3)
```

plotDSEA*Plotting functions for DSEA lists***Description**

Plotting functions for DSEA lists

Usage

```
plotDSEA(dsepso, dsesso, dsepi, dsepisem, dsfenics, dsspace, k)
```

Arguments

| | |
|----------|---|
| dsepso | list with enrichment for EpSO |
| dsesso | list with enrichment for ESSO |
| dsepi | list with enrichment for EPILONT |
| dsepisem | list with enrichment for EPISEM |
| dsfenics | list with enrichment for FENICS |
| dsspace | list with enrichment for the combined ranked list |
| k | numeric value for the length to be plotted |

Value

the plot object

Examples

```

utils::data(rawDrugNamesCo0cEpSO, package="epos")
utils::data(rawDrugNamesCo0cESSO, package="epos")
utils::data(rawDrugNamesCo0cEPILONT, package="epos")
utils::data(rawDrugNamesCo0cEPISEM, package="epos")
utils::data(rawDrugNamesCo0cFENICS, package="epos")
attrashda <-
  readAtcMapIntoHashMapDrugNamesAtcCodes(
    system.file("extdata", "db-atc.map", package = "epos"), "\t")
epso <- rawDrugNamesCo0cEpSO
neuroepso <- filterNeuroDrugs(epso, attrashda)
esso <- rawDrugNamesCo0cESSO
neuroesso <- filterNeuroDrugs(esso, attrashda)
epi <- rawDrugNamesCo0cEPILONT
neuroepi <- filterNeuroDrugs(epi, attrashda)
episem <- rawDrugNamesCo0cEPISEM
neuroepisem <- filterNeuroDrugs(episem, attrashda)
fenics <- rawDrugNamesCo0cFENICS
neurofenics <- filterNeuroDrugs(fenics, attrashda)
mx <- max(
  c(length(neuroepso), length(neuroesso), length(neuroepi),
    length(neuroepisem), length(neurofenics)))
dneuro <-
  data.frame(EpSO = c(neuroepso, rep("", (mx-length(neuroepso)))), 
             ESSO = c(neuroesso, rep("", (mx-length(neuroesso)))), 
             EPILONT = c(neuroepi, rep("", (mx-length(neuroepi)))), 
             EPISEM = c(neuroepisem, rep("", (mx-length(neuroepisem)))), 
             FENICS = c(neurofenics, rep("", (mx-length(neurofenics)))))
dneuromaxk <- TopKlists::calculate.maxK(dneuro, L=5, d=5, v=5)
neurospace <- as.character(dneuromaxk$topkspace)
dsepso <- calcDSEA(neuroepso, mx)
dsesso <- calcDSEA(neuroesso, mx)
dsepi <- calcDSEA(neuroepi, mx)
dsepisem <- calcDSEA(neuroepisem, mx)
dsfenics <- calcDSEA(neurofenics, mx)
dsspace <- calcDSEA(neurospace, mx)
p <- plotDSEA(dsepso, dsesso, dsepi, dsepisem, dsfenics, dsspace, dneuromaxk$maxK)
## Not run:
ggplot2::ggsave("dsea.png",
  p <- plotDSEA(dsepso, dsesso, dsepi, dsepisem, dsfenics, dsspace,
  dneuromaxk$maxK), width=480, height=320, units = "mm", dpi = 300)

## End(Not run)

```

Description

Plotting functions for enrichment lists

Usage

```
plotEnrichment(eposo, esso, epi, episem, fenics, space, k)
```

Arguments

| | |
|--------|---|
| eposo | list with enrichment for EpSO |
| esso | list with enrichment for ESSO |
| epi | list with enrichment for EPILONT |
| episem | list with enrichment for EPISEM |
| fenics | list with enrichment for FENICS |
| space | list with enrichment for the combined ranked list |
| k | numeric value for the length to be plotted |

Value

the plot object

Examples

```
utils::data(rawDrugNamesCo0cEpSO, package="epos")
utils::data(rawDrugNamesCo0cESSO, package="epos")
utils::data(rawDrugNamesCo0cEPILONT, package="epos")
utils::data(rawDrugNamesCo0cEPISEM, package="epos")
utils::data(rawDrugNamesCo0cFENICS, package="epos")
atHashda <-
  readAtcMapIntoHashMapDrugNamesAtcCodes(
    system.file("extdata", "db-atc.map", package = "epos"), "\t")
epso <- rawDrugNamesCo0cEpSO
neuroepso <- filterNeuroDrugs(epso, atHashda)
esso <- rawDrugNamesCo0cESSO
neuroesso <- filterNeuroDrugs(esso, atHashda)
epi <- rawDrugNamesCo0cEPILONT
neuroepi <- filterNeuroDrugs(epi, atHashda)
episem <- rawDrugNamesCo0cEPISEM
neuroepisem <- filterNeuroDrugs(episem, atHashda)
fenics <- rawDrugNamesCo0cFENICS
neurofenics <- filterNeuroDrugs(fenics, atHashda)
mx <- max(
  c(length(neuroepso), length(neuroesso), length(neuroepi),
    length(neuroepisem), length(neurofenics)))
dneuro <-
  data.frame(EpSO = c(neuroepso, rep("", (mx-length(neuroepso)))),
             ESSO = c(neuroesso, rep("", (mx-length(neuroesso)))),
             EPILONT = c(neuroepi, rep("", (mx-length(neuroepi)))),
             EPISEM = c(neuroepisem, rep("", (mx-length(neuroepisem)))),
```

```

FENICS = c(neurofenics, rep("", (mx-length(neurofenics)))))

dneuromaxk <- TopKLists::calculate.maxK(dneuro, L=5, d=5, v=5)
neurospace <- as.character(dneuromaxk$topkspace)
enepso <- calcEnrichment(neuroepso)
nesso <- calcEnrichment(neuroesso)
enepi <- calcEnrichment(neuroepi)
enepisem <- calcEnrichment(neuroepisem)
enfenics <- calcEnrichment(neurofenics)
enspace <- calcEnrichment(neurospace)
p <- plotEnrichment(enepso, nesso, enepi, enepisem, enfenics, enspace, dneuromaxk$maxK)

```

`printTop10Drugs`*Print Top 10 Drugs***Description**

Print Top 10 Drugs

Usage

```
printTop10Drugs(neuroepso, neuroesso, neuroepi, neuroepisem, neurofenics)
```

Arguments

| | |
|--------------------------|---|
| <code>neuroepso</code> | Ranked list of drug names co-occurring with EpSO |
| <code>neuroesso</code> | Ranked list of drug names co-occurring with ESSO |
| <code>neuroepi</code> | Ranked list of drug names co-occurring with EPILONT |
| <code>neuroepisem</code> | Ranked list of drug names co-occurring with EPISEM |
| <code>neurofenics</code> | Ranked list of drug names co-occurring with FENICS |

Value

data frame with top 10 drugs for each ontology

Examples

```

utils::data(rawDrugNamesCoOcEpSO, package="epos")
utils::data(rawDrugNamesCoOcESSO, package="epos")
utils::data(rawDrugNamesCoOcEPILONT, package="epos")
utils::data(rawDrugNamesCoOcEPISEM, package="epos")
utils::data(rawDrugNamesCoOcFENICS, package="epos")
atcHashda <-
readAtcMapIntoHashMapDrugNamesAtcCodes(
  system.file("extdata", "db-atc.map", package = "epos"), "\t")
atcHashaa <-
readAtcMapIntoHashMapAtcCodesAtcNames(
  system.file("extdata", "db-atc.map", package = "epos"), "\t")
atcHashsec <-

```

```

readSecondLevelATC(
  system.file("extdata", "atc-secondlevel.map", package = "epos"), "\t")
epso <- rawDrugNamesCoOcEpsO
neuroepso <- filterNeuroDrugs(epso, atchashda)
esso <- rawDrugNamesCoOcESSO
neuroesso <- filterNeuroDrugs(esso, atchashda)
epi <- rawDrugNamesCoOcEPILONT
neuroepi <- filterNeuroDrugs(epi, atchashda)
episem <- rawDrugNamesCoOcEPISEM
neuroepisem <- filterNeuroDrugs(episem, atchashda)
fenics <- rawDrugNamesCoOcFENICS
neurofenics <- filterNeuroDrugs(fenics, atchashda)
top10table <- printTop10Drugs(neuroepso, neuroesso, neuroepi, neuroepisem, neurofenics)
## Not run:
print(xtable::xtable(top10table, type = "latex"),
  file = "top10table.tex")

## End(Not run)

```

rawDrugNamesCoOcEPILONT

List drug terms with their frequency co-occurring with terms from the EPILONT ontology in publications since 2015 from the BioASQ 2020 corpus.

Description

List drug terms with their frequency co-occurring with terms from the EPILONT ontology in publications since 2015 from the BioASQ 2020 corpus.

Usage

```
rawDrugNamesCoOcEPILONT
```

Format

A named list of drug term frequencies

Source

The text mining workflows for data generation are described in Mueller, Bernd and Hagelstein, Alexandra (2016) <doi:10.4126/FRL01-006408558>, Mueller, Bernd et al. (2017) <doi:10.1007/978-3-319-58694-6_22>, and Mueller, Bernd and Rebholz-Schuhmann, Dietrich (2020) <doi:10.1007/978-3-030-43887-6_52>. The source data set for generating the data co-occurrence lists is the BioASQ 2020 corpus. The source ontology for the creation of the dictionary is the Epilepsy Ontology (EPILONT) from <https://bioportal.bioontology.org/ontologies/EPILONT>

Examples

```
utils::data(rawDrugNamesCoOcEPILONT, package="epos")
```

```
rawDrugNamesCoOcEPISEM
```

List drug terms with their frequency co-occurring with terms from the EPISEM ontology in publications since 2015 from the BioASQ 2020 corpus.

Description

List drug terms with their frequency co-occurring with terms from the EPISEM ontology in publications since 2015 from the BioASQ 2020 corpus.

Usage

```
rawDrugNamesCoOcEPISEM
```

Format

A named list of drug term frequencies

Source

The text mining workflows for data generation are described in Mueller, Bernd and Hagelstein, Alexandra (2016) <doi:10.4126/FRL01-006408558>, Mueller, Bernd et al. (2017) <doi:10.1007/978-3-319-58694-6_22>, and Mueller, Bernd and Rebholz-Schuhmann, Dietrich (2020) <doi:10.1007/978-3-030-43887-6_52>. The source data set for generating the data co-occurrence lists is the BioASQ 2020 corpus. The source ontology for the creation of the dictionary is the Epilepsy Semiology Ontology (EPISEM) from <https://bioportal.bioontology.org/ontologies/EPISEM>

Examples

```
utils::data(rawDrugNamesCoOcEPISEM, package="epos")
```

```
rawDrugNamesCoOcEpSO
```

List drug terms with their frequency co-occurring with terms from the EpSO ontology in publications since 2015 from the BioASQ 2020 corpus.

Description

List drug terms with their frequency co-occurring with terms from the EpSO ontology in publications since 2015 from the BioASQ 2020 corpus.

Usage

```
rawDrugNamesCoOcEpSO
```

Format

A named list of drug term frequencies

Source

The text mining workflows for data generation are described in Mueller, Bernd and Hagelstein, Alexandra (2016) <doi:10.4126/FRL01-006408558>, Mueller, Bernd et al. (2017) <doi:10.1007/978-3-319-58694-6_22>, and Mueller, Bernd and Rebholz-Schuhmann, Dietrich (2020) <doi:10.1007/978-3-030-43887-6_52>. The source data set for generating the data co-occurrence lists is the BioASQ 2020 corpus. The source ontology for the creation of the dictionary is the Epilepsy and Seizure Ontology (EpSO) from <https://bioportal.bioontology.org/ontologies/EPSO>

Examples

```
utils::data(rawDrugNamesCoOcEpSO, package="epos")
```

`rawDrugNamesCoOcESSO`

List drug terms with their frequency co-occurring with terms from the ESSO ontology in publications since 2015 from the BioASQ 2020 corpus.

Description

List drug terms with their frequency co-occurring with terms from the ESSO ontology in publications since 2015 from the BioASQ 2020 corpus.

Usage

```
rawDrugNamesCoOcESSO
```

Format

An object of class character of length 8620.

Source

The text mining workflows for data generation are described in Mueller, Bernd and Hagelstein, Alexandra (2016) <doi:10.4126/FRL01-006408558>, Mueller, Bernd et al. (2017) <doi:10.1007/978-3-319-58694-6_22>, and Mueller, Bernd and Rebholz-Schuhmann, Dietrich (2020) <doi:10.1007/978-3-030-43887-6_52>. The source data set for generating the data co-occurrence lists is the BioASQ 2020 corpus. The source ontology for the creation of the dictionary is Epilepsy Syndrome Seizure Ontology (ESSO) from <https://bioportal.bioontology.org/ontologies/ESSO>

Examples

```
utils::data(rawDrugNamesCoOcESSO, package="epos")
```

```
rawDrugNamesCoOcFENICS
```

List drug terms with their frequency co-occurring with terms from the FENICS ontology in publications from the BioASQ 2020 corpus.

Description

List drug terms with their frequency co-occurring with terms from the FENICS ontology in publications from the BioASQ 2020 corpus.

Usage

```
rawDrugNamesCoOcFENICS
```

Format

A named list of drug term frequencies

Source

The text mining workflows for data generation are described in Mueller, Bernd and Hagelstein, Alexandra (2016) <doi:10.4126/FRL01-006408558>, Mueller, Bernd et al. (2017) <doi:10.1007/978-3-319-58694-6_22>, and Mueller, Bernd and Rebholz-Schuhmann, Dietrich (2020) <doi:10.1007/978-3-030-43887-6_52>. The source data set for generating the data co-occurrence lists is the BioASQ 2020 corpus. The source ontology for the creation of the dictionary is the Functional Epilepsy Nomenclature for Ion Channels (FENICS) from <https://bioportal.bioontology.org/ontologies/FENICS>

Examples

```
utils::data(rawDrugNamesCoOcFENICS, package="epos")
```

```
readAtcMapIntoHashMapAtcCodesAtcNames
```

Processes the input file db-atc.map to form a HashMap containing the drug names with ATC codes

Description

Processes the input file db-atc.map to form a HashMap containing the drug names with ATC codes

Usage

```
readAtcMapIntoHashMapAtcCodesAtcNames(filename, separator)
```

Arguments

| | |
|------------------------|--|
| <code>filename</code> | character vector with the file name of the file db-atc.map |
| <code>seperator</code> | character vector with the seperator used within the map-file |

Value

atcHashaa hash with atc codes as keys and atc names as values

Examples

```
atcHashaa <-
  readAtcMapIntoHashMapAtcCodesAtcNames(
    system.file("extdata", "db-atc.map", package = "epos"), "\t")
```

`readAtcMapIntoHashMapDrugNamesAtcCodes`

Processes the input file db-atc.map to form a HashMap containing the drug names with ATC codes

Description

Processes the input file db-atc.map to form a HashMap containing the drug names with ATC codes

Usage

```
readAtcMapIntoHashMapDrugNamesAtcCodes(filename, seperator)
```

Arguments

| | |
|------------------------|--|
| <code>filename</code> | character vector with the file name of the file db-atc.map |
| <code>seperator</code> | character vector with the seperator used within the map-file |

Value

atcHashda hash with drug names as keys and atc codes as values

Examples

```
atcHashda <- readAtcMapIntoHashMapDrugNamesAtcCodes(
  system.file("extdata", "db-atc.map", package = "epos"), "\t")
```

readSecondLevelATC *Read the second level ATC classes from the file atc-secondlevel.map*

Description

Read the second level ATC classes from the file atc-secondlevel.map

Usage

```
readSecondLevelATC(filename, separator)
```

Arguments

| | |
|-----------|--|
| filename | the file name that is supposed to be atc-secondlevel.map |
| separator | the csv file delimiter |

Value

atcHashSec a hash with second level ATC classes as keys and their names as values

Examples

```
atcHashSec <-  
  readSecondLevelATC(  
    system.file("extdata", "atc-secondlevel.map", package = "epos"), "\t")
```

sortTableByRefMatches *Sort table by scoring for each row*

Description

Sort table by scoring for each row

Usage

```
sortTableByRefMatches(dntk)
```

Arguments

| | |
|------|---|
| dntk | the table returned from writeNeuroTable |
|------|---|

Value

the sorted table

Examples

```

utils::data(rawDrugNamesCoOcEpSO, package="epos")
utils::data(rawDrugNamesCoOcESSO, package="epos")
utils::data(rawDrugNamesCoOcEPILONT, package="epos")
utils::data(rawDrugNamesCoOcEPISEM, package="epos")
utils::data(rawDrugNamesCoOcFENICS, package="epos")
atcHashda <-
readAtcMapIntoHashMapDrugNamesAtcCodes(
  system.file("extdata", "db-atc.map", package = "epos"), "\t")
atcHashaa <-
readAtcMapIntoHashMapAtcCodesAtcNames(
  system.file("extdata", "db-atc.map", package = "epos"), "\t")
atcHashsec <-
readSecondLevelATC(
  system.file("extdata", "atc-secondlevel.map", package = "epos"), "\t")
epso <- rawDrugNamesCoOcEpSO
neuroepso <- filterNeuroDrugs(epso, atcHashda)
esso <- rawDrugNamesCoOcESSO
neuroesso <- filterNeuroDrugs(esso, atcHashda)
epi <- rawDrugNamesCoOcEPILONT
neuroepi <- filterNeuroDrugs(epi, atcHashda)
episem <- rawDrugNamesCoOcEPISEM
neuroepisem <- filterNeuroDrugs(episem, atcHashda)
fenics <- rawDrugNamesCoOcFENICS
neurofenics <- filterNeuroDrugs(fenics, atcHashda)
mx <- max(
  c(length(neuroepso), length(neuroesso), length(neuroepi),
    length(neuroepisem), length(neurofenics)))
dneuro <-
data.frame(EpSO = c(neuroepso, rep("", (mx-length(neuroepso)))),
           ESSO = c(neuroesso, rep("", (mx-length(neuroesso)))),
           EPILONT = c(neuroepi, rep("", (mx-length(neuroepi)))),
           EPISEM = c(neuroepisem, rep("", (mx-length(neuroepisem)))),
           FENICS = c(neurofenics, rep("", (mx-length(neurofenics)))))
suppressWarnings(dneuromaxk <- TopKLists::calculate.maxK(dneuro, L=5, d=5, v=5))
neurotable <- createNeuroTable(atcHashda, atcHashsec, dneuromaxk)
sortedNeuroTable <- sortTableByRefMatches(neurotable)
## Not run:
print(xtable::xtable(sortedNeuroTable, type = "latex"),
      file = "sortedNeuroTable.tex",
      include.rownames=FALSE)

## End(Not run)

```

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