Package 'UniprotR'

July 3, 2022

Title Retrieving Information of Proteins from Uniprot
Version 2.2.0
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Description Connect to Uniprot https://www.uniprot.org/ to retrieve information about proteins using their accession number such information could be name or taxonomy information, For detailed information kindly read the publication https://www.sciencedirect.com/science/article/pii/S1874391919303859 .
License GPL-3
Encoding UTF-8
RoxygenNote 7.1.2
Imports utils, grDevices, stats, grid, graphics, httr, plyr, dplyr, scales, magrittr, magick, data.tree, ggplot2, tidyverse, gridExtra, ggpubr, curl, networkD3, stringr, qdapRegex, htmlwidgets, alakazam (>= 1.0.0), gprofiler2, progress, ggsci
<pre>URL https://github.com/Proteomicslab57357/UniprotR</pre>
<pre>BugReports https://github.com/Proteomicslab57357/UniprotR/issues</pre>
NeedsCompilation no
Repository CRAN
Date/Publication 2022-07-03 12:30:08 UTC
R topics documented:
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Description

This Function is used to plot Genes Tree in the data of the accession/s.

Usage

ConstructGenesTree(ProteinDataObject,directorypath = NULL)

Arguments

ProteinDataObject

input a Dataframe returned from GetNamesTaxa function directorypath path to save txt file containing results returened by the function.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

ConstructLocTree Connect and parse UniProt information.

Description

This Function is used to plot location's Tree in the data of the accession/s in the chromosomes.

Usage

ConstructLocTree(ProteinDataObject, directorypath = NULL)

Arguments

ProteinDataObject

input a Dataframe returned from GetNamesTaxa function directorypath path to save txt file containing results returened by the function.

Author(s)

4 Enrichment.BP

ConvertID	Connect and parse UniProt database identifiers information.

Description

The function is work to convert the UniProtKB AC/ID to any database identifiers available by the UniProtKB. For more information about available database identifiers see https://www.uniprot.org/help/api_idmapping.

Usage

```
ConvertID(ProteinAccList , ID_from = "ACC+ID" , ID_to = NULL
  , directorypath = NULL)
```

Arguments

ProteinAccList Vector of UniProt Accession/s

ID_from string of database identifier abbreviation, from which the Accession/ID will be

converted

ID_to string of database identifier abbreviation, to which the Accession/ID will be

converted. default is all database identifier available in UniProtKB

directorypath path to save excel file containing results returened by the function.

Value

DataFrame where column one contains the Accession/ID before conversion and other columns contains the Accession/ID after conversion

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

|--|

Description

This function is used for Enrichment analysis of biological process of given list of genes or proteins

Usage

```
Enrichment.BP(Accs,OS="hsapiens",p_value=0.05,directorypath=NULL)
```

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Arguments

Accs Vector of UniProt Accession/s or genes

OS organism name Example: human - 'hsapiens', mouse - 'mmusculus'

p_value custom p-value threshold for significance, default = 0.05

directorypath Path to save output plot

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Enrichment.CC Connect and parse UniProt information

Description

This function is used for Enrichment analysis of cellular component of given list of genes or proteins

Usage

```
Enrichment.CC(Accs,OS="hsapiens",p_value=0.05,directorypath=NULL)
```

Arguments

Accs Vector of UniProt Accession/s or genes

OS organism name Example: human - 'hsapiens', mouse - 'mmusculus'

 p_{value} custom p-value threshold for significance, default = 0.05

directorypath Path to save output plot

Author(s)

6 Enrichment.MF

Enrichment.KEGG Connect and parse UniProt information

Description

This function is used for Enrichment analysis of given list of genes or proteins from KEGG database

Usage

```
Enrichment.KEGG(Accs,OS="hsapiens",p_value=0.05,directorypath=NULL)
```

Arguments

Accs Vector of UniProt Accession/s or genes

OS organism name Example: human - 'hsapiens', mouse - 'mmusculus'

p_value custom p-value threshold for significance, default = 0.05

directorypath Path to save output plot

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

|--|--|

Description

This function is used for Enrichment analysis of Molecular function of given list of genes or proteins

Usage

```
Enrichment.MF(Accs,OS="hsapiens",p_value=0.05,directorypath=NULL)
```

Arguments

Accs Vector of UniProt Accession/s or genes

OS organism name Example: human - 'hsapiens', mouse - 'mmusculus'

p_value custom p-value threshold for significance, default = 0.05

directorypath Path to save output plot

Author(s)

Enrichment.REAC 7

Enrichment.REAC	Connect and parse UniProt information	

Description

This function is used for Enrichment analysis of given list of genes or proteins from REACTOME

Usage

```
Enrichment.REAC(Accs,OS="hsapiens",p_value=0.05,directorypath=NULL)
```

Arguments

Accs Vector of UniProt Accession/s or genes

OS organism name Example: human - 'hsapiens', mouse - 'mmusculus'

p_value custom p-value threshold for significance, default = 0.05

directorypath Path to save output plot

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Get.diseases Connect

Connect and parse UniProt information.

Description

This Function is used to get diseases associated with proteins.

Usage

```
Get.diseases(Pathology_object , directorypath = NULL)
```

Arguments

Pathology_object

Dataframe retrieved from UniprotR Function "GetPathology_Biotech"

directorypath path to save Output file

Author(s)

8 GetExpression

GetAccessionList

Connect and parse UniProt information.

Description

This function can be used to get a list of UniProt Accession/s from a csv file.

Usage

```
GetAccessionList(DataObjPath)
```

Arguments

DataObjPath input path of excel file

Value

a vector of UniProt Accession/s

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

GetExpression

Connect and parse UniProt Expression information.

Description

The function is work to retrieve Expression data from UniProt for a list of proteins accessions.For more information about what included in the Expression data see https://www.uniprot.org/help/return_fields.

Usage

```
GetExpression(ProteinAccList , directorypath = NULL)
```

Arguments

ProteinAccList Vector of UniProt Accession/s

directorypath path to save excel file containing results returened by the function.

Value

DataFrame where rows names are the accession and columns contains the information retrieved from the UniProt

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Note

The function also, Creates a csv file with the retrieved information.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

GetFamily_Domains

Connect and parse UniProt Family Domains information.

Description

The function is work to retrieve Family Domains data from UniProt for a list of proteins accessions. For more information about what included in the Family Domains data see https://www.uniprot.org/help/return_fields.

Usage

```
GetFamily_Domains(ProteinAccList , directorypath = NULL)
```

Arguments

ProteinAccList Vector of UniProt Accession/s

directorypath path to save excel file containing results returened by the function.

Value

DataFrame where rows names are the accession and columns contains the information retrieved from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

GetGeneral_Information

Connect and parse UniProt General Information.

Description

The function is work to retrieve General Information data from UniProt for a list of proteins accessions. For more information about what included in the General Information data see https://www.uniprot.org/help/return_field

Usage

```
GetGeneral_Information(ProteinAccList , directorypath = NULL)
```

Arguments

ProteinAccList Vector of UniProt Accession/s
directorypath path to save excel file containing results returened by the function.

Value

DataFrame where rows names are the accession and columns contains the General Information of protein from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

```
Obj <- GetGeneral_Information("014520")</pre>
```

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GetMiscellaneous	Connect and parse UniProt Miscellaneous information.
Getiliscerraneous	Connect and purse Only for Miscentineous information.

Description

The function is work to retrieve Miscellaneous data from UniProt for a list of proteins accessions. For more information about what included in the Miscellaneous data see https://www.uniprot.org/help/return_fields.

Usage

```
GetMiscellaneous(ProteinAccList , directorypath = NULL)
```

Arguments

```
ProteinAccList Vector of UniProt Accession/s
directorypath path to save excel file containing results returened by the function.
```

Value

DataFrame where rows names are the accession and columns contains the information retrieved from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

```
Obj <- GetMiscellaneous("014520")
```

GetNamesTaxa	Connect and parse UniProt Names Taxa information.	

Description

The function is work to retrieve Names Taxa data from UniProt for a list of proteins accessions. For more information about what included in the NamesTaxa data see https://www.uniprot.org/help/return_fields.

Usage

```
GetNamesTaxa(ProteinAccList , directorypath = NULL)
```

Arguments

```
ProteinAccList Vector of UniProt Accession/s
directorypath path to save excel file containing results returened by the function.
```

Value

DataFrame where rows names are the accession and columns contains the information of protein name & taxonomy from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

```
Obj <- GetNamesTaxa("014520")
```

GetPathology_Biotech Connect and parse UniProt Pathology_Biotech information.

Description

The function is work to retrieve Pathology_Biotech data from UniProt for a list of proteins accessions. For more information about what included in the Pathology_Biotech data see https://www.uniprot.org/help/return_fields

Usage

```
GetPathology_Biotech(ProteinAccList , directorypath = NULL)
```

Arguments

```
ProteinAccList Vector of UniProt Accession/s
directorypath path to save excel file containing results returened by the function.
```

Value

DataFrame where rows names are the accession and columns contains the information retrieved from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

GetpdbStructure 13

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

```
Obj <- GetPathology_Biotech("014520")</pre>
```

GetpdbStructure

Connect and parse UniProt information.

Description

The function is work to retrieving GetpdbStructure and download it to user directory.

Usage

```
GetpdbStructure(ProteinAccList , directorypath = NULL)
```

Arguments

```
ProteinAccList input a vector of UniProt Accession/s directorypath path to save excel file containing results returened by the function.
```

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

GetProteinAnnontate

Connect and parse UniProt information.

Description

The function is work to retrieve user-defined information data from UniProt for a list of proteins accessions For more information see https://www.uniprot.org/help/uniprotkb_column_names

Usage

```
GetProteinAnnontate(ProteinAccList , columns)
```

Arguments

```
ProteinAccList a vector of UniProt Accession/s columns a vector of UniProtKB column names
```

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Value

DataFrame where rows names are the accession and columns contains the information retrieved from the UniProt

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

```
Obj <- GetProteinInteractions("014520")</pre>
```

GetProteinFunction

Connect and parse UniProt Protein Function information.

Description

The function is work to retrieve Protein Function data from UniProt for a list of proteins accessions. For more information about what included in the Protein Function data see https://www.uniprot.org/help/return_fields.

Usage

```
GetProteinFunction(ProteinAccList , directorypath = NULL)
```

Arguments

```
ProteinAccList Vector of UniProt Accession/s
directorypath path to save excel file containing results returened by the function.
```

Value

DataFrame where rows names are the accession and columns contains the information of protein function roles from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

```
Obj <- GetProteinFunction("014520")</pre>
```

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GetProteinGOInfo

Connect and parse UniProt proteins gene ontology information.

Description

The function is work to retrieve proteins gene ontology data from UniProt for a list of proteins accessions. For more information about what included in the proteins gene ontology data see https://www.uniprot.org/help/return_fi

Usage

```
GetProteinGOInfo(ProteinAccList , directorypath = NULL)
```

Arguments

```
ProteinAccList Vector of UniProt Accession/s
directorypath path to save excel file containing results returened by the function.
```

Value

DataFrame where rows names are the accession and columns contains the information of Gene ontology of protein from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

```
Obj <- GetProteinGOInfo("014520")</pre>
```

GetProteinInteractions

Connect and parse UniProt protein Interactions information.

Description

The function is work to retrieve protein Interactions data from UniProt for a list of proteins accessions. For more information about what included in the protein Interactions data see https://www.uniprot.org/help/return_fields

Usage

```
GetProteinInteractions(ProteinAccList , directorypath = NULL)
```

16 GetproteinNetwork

Arguments

```
ProteinAccList Vector of UniProt Accession/s
directorypath path to save excel file containing results returened by the function.
```

Value

DataFrame where rows names are the accession and columns contains the information of protein Interactions from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

```
Obj <- GetProteinInteractions("ProteinAccession")</pre>
```

GetproteinNetwork

Connect and parse stringdb information.

Description

This function is connecting to stringdb and retrieve all possible interactions for the searched protein/s.

Usage

```
GetproteinNetwork(ProteinAccList , directorypath = NULL)
GetproteinNetwork(ProteinAccList , directorypath = NULL)
```

Arguments

```
ProteinAccList input a vector of UniProt Accession/s
directorypath path to save excel file containing results returened by the function.
```

Author(s)

GetproteinNetwork_all

GetproteinNetwork_all Connect and parse stringdb information.

Description

This function is connecting to stringdb and retrieve PPI between input list

Usage

```
GetproteinNetwork_all(ProteinAccList , directorypath = NULL, SpeciesID = 9606)
GetproteinNetwork_all(ProteinAccList , directorypath = NULL, SpeciesID = 9606)
```

Arguments

ProteinAccList input a vector of UniProt Accession/s

directorypath path to save excel file containing results returened by the function.

SpeciesID Taxonomic id of accession's species ex. homo sapines 9606

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

GetProteomeFasta

Connect and DOWNLOAD Proteome info.

Description

The function is work to retrieve proteome information in FASTA format based on proteome id.

Usage

```
GetProteomeFasta(ProteomeID , directorypath = NULL)
```

Arguments

Proteome ID from UniProt

directorypath path to save FASTA file containing results returened by the function.

Note

The function Download fasta format of proteome.

Author(s)

GetProteomeInfo

Connect and DOWNLOAD Proteome info.

Description

The function is work to retrieve proteome information based on proteome id.

Usage

```
GetProteomeInfo(ProteomeID , directorypath = NULL)
```

Arguments

Proteome ID from UniProt

directorypath path to save CSV file containing results returened by the function.

Note

The function Download csv Info of proteome.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

GetPTM_Processing

Connect and parse UniProt PTM_Processing information.

Description

The function is work to retrieve PTM_Processing data from UniProt for a list of proteins accessions. For more information about what included in the PTM_Processing data see https://www.uniprot.org/help/return_fields

Usage

```
GetPTM_Processing(ProteinAccList, directorypath = NULL)
```

Arguments

ProteinAccList Vector of UniProt Accession/s

directorypath path to save excel file containing results returened by the function

Value

DataFrame where rows names are the accession and columns contains the information retrieved from the UniProt

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Note

The function also, Creates a csv file with the retrieved information.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

```
Obj <- GetPTM_Processing("014520" )</pre>
```

GetPublication

Connect and parse UniProt Publication about a protein information.

Description

The function is work to retrieve Publication data from UniProt for a list of proteins accessions. For more information about what included in the Publication data see https://www.uniprot.org/help/return_fields.

Usage

```
GetPublication(ProteinAccList , directorypath = NULL)
```

Arguments

```
ProteinAccList Vector of UniProt Accession/s
directorypath path to save excel file containing results returened by the function.
```

Value

DataFrame where rows names are the accession and columns contains the Publication of protein from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

20 GetSeqLength

GETSeqFastaUniprot	Connect and parse UniProt information.
--------------------	--

Description

This Function is used to get Sequence information of accession/s from Uniprot as a Fasta file.

Usage

```
GETSeqFastaUniprot(Accessions,FilePath = NULL, FileName = NULL)
```

Arguments

Accessions Vector of UniProt Accession/s

FilePath path of directory to save the output fasta.

FileName Name of the fasta file.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

GetSeqLength	Connect and parse UniProt Sequences information.

Description

The function is work to retrieve Sequence's Length data from Uniparc for a list of proteins accessions. This function was added to overcome the NAs returned when Uniprot database deleted the protein from the database

Usage

```
GetSeqLength(ProteinAccList, directorypath = NULL)
```

Arguments

```
ProteinAccList Vector of UniProt Accession/s
directorypath path to save excel file containing results returened by the function
```

Value

DataFrame where rows names are the accession and columns contains the information retrieved from the UniProt

GetSequenceIso 21

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

```
Obj <- GetSeqLength("014520")</pre>
```

GetSequenceIso

Connect and parse UniProt Sequences information.

Description

The function is work to retrieve protein's Sequence data from Uniparc for a list of proteins accessions. This function was added to handle isoformes

Usage

```
GetSequenceIso(ProteinAccList, directorypath = NULL)
```

Arguments

ProteinAccList Vector of UniProt Accession/s
directorypath path to save excel file containing results returened by the function

Value

DataFrame where rows names are the accession and columns contains the information retrieved from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

```
Obj <- GetSequenceIso("014520")</pre>
```

22 GetStructureInfo

GetSequences

Connect and parse UniProt Sequences information.

Description

The function is work to retrieve Sequences data from UniProt for a list of proteins accessions. For more information about what included in the Sequences data see https://www.uniprot.org/help/return_fields.

Usage

```
GetSequences(ProteinAccList, directorypath = NULL)
```

Arguments

```
ProteinAccList Vector of UniProt Accession/s
directorypath path to save excel file containing results returened by the function
```

Value

DataFrame where rows names are the accession and columns contains the information retrieved from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

```
Obj <- GetSequences("014520")</pre>
```

GetStructureInfo

Connect and parse UniProt protein Structure information.

Description

The function is work to retrieve Structral data from UniProt for a list of proteins accessions. For more information about what included in the structral data see https://www.uniprot.org/help/return_fields.

Usage

```
GetStructureInfo(ProteinAccList, directorypath = NULL)
```

GetSubcellular_location

Arguments

ProteinAccList Vector of UniProt Accession/s.

directorypath path to save excel file containing results returened by the function.

Value

DataFrame where rows names are the accession and columns contains the Structural information of protein from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

GetSubcellular_location

Connect and parse UniProt protein Subcellular location information.

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Description

The function is work to retrieve protein Subcellular location data from UniProt for a list of proteins accessions. For more information about what included in the protein Subcellular location data see https://www.uniprot.org/help/return_fields.

Usage

GetSubcellular_location(ProteinAccList, directorypath = NULL)

Arguments

ProteinAccList Vector of UniProt Accession/s.

directorypath path to save excel file containig results returened by the function.

Value

DataFrame where rows names are the accession and columns contains the information about Subcellular location of protein from the UniProt

Note

The function also, Creates a csv file with the retrieved information.

Author(s)

24 HandleBadRequests

Goparse

Connect and parse UniProt information.

Description

This Function is used to parse data retrieved from UniprotR Function "GetProteinGOInfo".

Usage

```
Goparse(GOObj, index = 3)
```

Arguments

G00bj Dataframe.

index idex of Go term in GoObj

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

HandleBadRequests

Handle bad requests This Function is used to handle possible errors when trying to get url response.

Description

Handle bad requests This Function is used to handle possible errors when trying to get url response.

Usage

HandleBadRequests(RequestCode)

Arguments

RequestCode Response re

Response returned from url

Value

None

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Examples

HandleBadRequests(400)

Pathway.Enr 25

Pathway.Enr Connect and parse UniProt information

Description

This function is used for Enrichment analysis of given list of genes or proteins

Usage

```
Pathway.Enr(Accs,OS="hsapiens",p_value=0.05,directorypath=NULL)
```

Arguments

Accs Vector of UniProt Accession/s or genes

OS organism Example: human - 'hsapiens' for more info https://biit.cs.ut.ee/gprofiler/page/organism-

list

p_value custom p-value threshold for significance, default = 0.05

directorypath Path to save output plot

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

Plot.GOMolecular	Connect and parse UniProt information.

Description

This Function is used to plot Molecular function of proteins.

Usage

```
Plot.GOMolecular(GOObj, Top = 10, directorypath = NULL)
```

Arguments

GOObj Dataframe returned from UniprotR Function "GetProteinGOInfo"

Top Number of molecular functions to be visualized

directorypath path to save Output plot.

Author(s)

26 PlotAcidity

Plot.GOSubCellular

Connect and parse UniProt information.

Description

This Function is used to plot subcellular localization of proteins.

Usage

```
Plot.GOSubCellular(GOObj, Top = 10, directorypath = NULL)
```

Arguments

G00bj Dataframe returned from UniprotR Function "GetProteinGOInfo"

Top Number of molecular functions to be visualized

directorypath path to save Output plot.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

PlotAcidity

Connect and parse UniProt information.

Description

This Function is used to plot proteins acidity retrieved from "GetSequences" Function.

Usage

```
PlotAcidity(SeqDataObjPath , directorypath = NULL)
```

Arguments

```
SeqDataObjPath Dataframe retrieved from UniprotR Function "GetSequences" directorypath path to save Output plot.
```

Author(s)

PlotCharge 27

PlotCharge

Connect and parse UniProt information.

Description

This Function is used to plot proteins charge retrieved from "GetSequences" Function.

Usage

```
PlotCharge(SeqDataObjPath , directorypath = NULL)
```

Arguments

```
SeqDataObjPath Dataframe retrieved from UniprotR Function "GetSequences" directorypath path to save Output plot.
```

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

PlotChromosomeInfo

Connect and parse UniProt information.

Description

This Function is used to plot location's frequency in the data of the accession/s in the chromosomes.

Usage

```
PlotChromosomeInfo(ProteinDataObject, directorypath = NULL)
```

Arguments

```
{\tt ProteinDataObject}
```

input a Dataframe returned from GetNamesTaxa function

directorypath path to save files returened by the function.

Author(s)

ot information	Connect and parse	PlotEnrichedGO
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Description

This function is used to generate a combined plot for the enriched Gene Ontology terms

Usage

PlotEnrichedGO(Accs, OS="hsapiens", p_value=0.05, Path=NULL, theme="aaas", width=7, height=7)

Arguments

Accs	Vector of UniProt Accession/s or genes
------	--

OS organism name Example: human - 'hsapiens', mouse - 'mmusculus'

p_value custom p-value threshold for significance, default = 0.05

Path to save output plot

theme optional parameter to generate specific theme for journals ex: "aaas", "nature",

"lancet", "jama"

width width of the generated plot height height of the generated plot

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

PlotEnrichedPathways Connect and parse UniProt information

Description

This function is used to generate a combined plot for the enriched pathways from KEGG and RE-ACTOME

Usage

PlotEnrichedPathways(Accs, OS="hsapiens", p_value=0.05, Path=NULL, theme="aaas", w=w, h=h)

PlotGenesNetwork 29

Arguments

Accs Vector of UniProt Accession/s or genes

OS organism name Example: human - 'hsapiens', mouse - 'mmusculus'

p_value custom p-value threshold for significance, default = 0.05

Path Path to save output plot

theme optional parameter to generate specific theme for journals ex: "aaas", "nature",

"lancet", "jama"

w width of the generated ploth height of the generated plot

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

PlotGenesNetwork Connect and parse UniProt information.

Description

This Function is used to cluster proteins based on primary genes retrieved from "GetNamesTaxa" Function.

Usage

PlotGenesNetwork(ProteinDataObject , directorypath = NULL)

Arguments

ProteinDataObject

Dataframe retrieved from UniprotR Function "GetNamesTaxa"

directorypath path to save Output plot.

Author(s)

30 PlotGOBiological

PlotGOAll	Connect and parse UniProt information.
	1

Description

This Function is used to plot the retrieved Gene Ontology from function 'GetProteinGOInfo'.

Usage

```
PlotGOAll(GOObj, Top = 10, directorypath = NULL, width = width, height = height)
```

Arguments

GOObj Dataframe returned from UniprotR Function "GetProteinGOInfo"

Top Number of molecular functions to be visualized

directorypath path to save Output plot.
width width of the generated plot
height height of the generated plot

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

PlotGOBiological Connect and parse UniProt information.

Description

This Function is used to plot Biological process of proteins.

Usage

```
PlotGOBiological(GOObj, Top = 10, directorypath = NULL)
```

Arguments

GOObj Dataframe returned from UniprotR Function "GetProteinGOInfo"

Top Number of molecular functions to be visualized

directorypath path to save Output plot.

Author(s)

PlotGoInfo 31

PΊ	otGoI	'nfn

Connect and parse UniProt information.

Description

This Function is used to plot data retrieved from UniprotR Function "GetProteinGOInfo".

Usage

```
PlotGoInfo(GOObj , directorypath = NULL)
```

Arguments

G00bj Dataframe retrieved from UniprotR Function "GetProteinGOInfo". directorypath path to save excel file containig results returened by the function (default = NA).

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

LT0	tGoter	ms

Connect and parse UniProt information.

Description

This Function is used to plot data retrieved from UniprotR Function "GetProteinGOInfo".

Usage

```
PlotGoterms(GOObj , directorypath = NULL)
```

Arguments

```
G00bj Dataframe.
```

directorypath path to save plot returned by function (default = NA).

Author(s)

32 PlotPhysicochemical

PlotGravy

Connect and parse UniProt information.

Description

This Function is used to plot proteins gravy index retrieved from "GetSequences" Function.

Usage

```
PlotGravy(SeqDataObjPath , directorypath = NULL)
```

Arguments

```
SeqDataObjPath Dataframe retrieved from UniprotR Function "GetSequences" directorypath path to save Output plot.
```

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

PlotPhysicochemical

Connect and parse UniProt information.

Description

This function can be used to get a list of UniProt Accession/s from a csv file.

Usage

```
PlotPhysicochemical(SeqDataObjPath , directorypath = NULL)
```

Arguments

```
SeqDataObjPath Dataframe returned from GetSequence function.
directorypath Path to save Physcochemical properties plot.
```

Author(s)

PlotproteinExist 33

PlotproteinExist

Connect and parse UniProt information.

Description

This Function is used to plot protein status in the data of the accession/s.

Usage

```
PlotproteinExist(ProteinDataObject,directorypath = NULL)
```

Arguments

ProteinDataObject

input a Dataframe returned from GetMiscellaneous function

directorypath path to save files returened by the function.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

PlotProteinGO_bio

Connect and parse UniProt information.

Description

This Function is used to plot biological process data retrieved from UniprotR Function "GetProtein-GOInfo".

Usage

```
PlotProteinGO_bio(GO_df , dir_path = NA)
```

Arguments

GO_df

Dataframe.

dir_path

path to save files returened by the function (default = NA).

Note

if no dir_path was given (default = NA) the function will only view the plot and will not save it

Author(s)

34 PlotProteinGO_molc

PlotProteinGO_cel

Connect and parse UniProt information.

Description

This Function is used to plot cellular components data retrieved from UniprotR Function "GetProteinGOInfo".

Usage

```
PlotProteinGO_cel(GO_df , dir_path = NA)
```

Arguments

GO_df Dataframe.

 dir_path path to save files returened by the function (default = NA).

Note

if no dir_path was given (default = NA) the function will only view the plot and Will not save it

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

PlotProteinGO_molc

Connect and parse UniProt information.

Description

This Function is used to plot molecular functions data retrieved from UniprotR Function "GetProteinGOInfo".

Usage

```
PlotProteinGO_molc(GO_df , dir_path = NA)
```

Arguments

GO_df Dataframe.

 dir_{path} path to save files returened by the function (default = NA).

Note

if no dir_path was given (default = NA) the function will only view the plot and will not save it

PlotproteinStatus 35

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

PlotproteinStatus

Connect and parse UniProt information.

Description

This Function is used to plot protein status in the data of the accession/s.

Usage

```
PlotproteinStatus (ProteinDataObject, directorypath = NULL)
```

Arguments

ProteinDataObject

input a Dataframe returned from GetMiscellaneous function

directorypath path to save files returened by the function.

Author(s)

Mohmed Soudy <Mohamed.soudy@57357.com> and Ali Mostafa <ali.mo.anwar@std.agr.cu.edu.eg>

PlotProteinTaxa

Connect and parse UniProt information.

Description

This Function is used to plot different taxas found of the accessions.

Usage

```
PlotProteinTaxa(ProteinDataObject , directorypath = NULL)
```

Arguments

ProteinDataObject

input a Dataframe of proteins as rownames.

directorypath path to save files returened by the function.

Author(s)

36 ProcessAcc

ProcessAcc

Connect and parse UniProt information.

Description

This Function is used to check validty of input accessions the data of the accession/s.

Usage

ProcessAcc(Accessions)

Arguments

Accessions

acession list returened from GetAccession function

Author(s)

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