Package 'biocViews'

February 20, 2025

Title Categorized views of R package repositories

Description Infrastructure to support 'views' used to classify Bioconductor packages. 'biocViews' are directed acyclic graphs of terms from a controlled vocabulary. There are three major classifications, corresponding to 'software', 'annotation', and 'experiment data' packages.

biocViews Infrastructure

URL http://bioconductor.org/packages/biocViews

BugReports https://github.com/Bioconductor/biocViews/issues

Version 1.75.0

License Artistic-2.0

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Maintainer Bioconductor Package Maintainer <maintainer@bioconductor.org>

Depends R (>= 3.6.0)

Imports Biobase, graph (>= 1.9.26), methods, RBGL (>= 1.13.5), tools, utils, XML, RCurl, RUnit, BiocManager

Suggests BiocGenerics, knitr, commonmark, BiocStyle

Collate AllClasses.R AllGenerics.R as-methods.R htmlDoc-methods.R htmlFilename-methods.R htmlValue-methods.R show-methods.R getPackNames.R packageDetails.R pump.R repository.R showvoc.R getPackageNEWS.R validation_tests.R recommendBiocViews.R dump_concept.R

VignetteBuilder knitr

git_url https://git.bioconductor.org/packages/biocViews

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biocViews-package

Description

Structures for vocabularies and narratives of views. This can be used to create HTML views of the package structure in a Bioconductor repository.

Details

| Package: | biocViews |
|------------|---|
| Version: | 1.11.4 |
| Depends: | R ($\geq 2.4.0$), methods, utils |
| Imports: | tools, Biobase, graph (>= 1.9.26), RBGL (>= 1.13.5), XML |
| Suggests: | Biobase |
| License: | Artistic-2.0 |
| URL: | http://www.bioconductor.org/packages/release/BiocViews.html |
| biocViews: | Infrastructure |

Index:

| BiocView-class Htmlized-class PackageDetail-class RepositoryDetail-class | Class "BiocView" Class "Htmlized" Class "PackageDetail" |
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| | Class "RepositoryDetail" |
| biocViewsVocab | Bioconductor Task Views Vocabulary Data |
| extractVignettes | Extract pdf vignettes from local package repository |
| genReposControlFiles | Generate CRAN-style repository control files |
| getBiocSubViews | Build a list of BiocView objects from a package repository |
| getBiocViews | Build a list of BiocView objects from a package repository |
| getPacksAndViews | Parse VIEWS file for views and packages |
| getSubTerms | Retrieve a term and its children from a vocab DAG |
| htmlDoc | Create a complete HTML document representation of an object |
| htmlFilename | Return a filename for an object's HTML representation |
| htmlValue | HTML Representation of an Object |
| writeBiocViews | Write a list of BiocView objects to HTML |
| writeHtmlDoc writePackageDetailHtml | Write an XML DOM containing HTML to a file |

| | Write HTML files for packages in a CRAN-style repository |
|---------------------|--|
| writeRepositoryHtml | Write package descriptions and a repository index as HTML |
| writeTopLevelView | Write the view for the root of a vocabulary to disk |
| write_REPOSITORY | Write a REPOSITORY control file for a CRAN-style package repository |
| write_SYMBOLS | Write a SYMBOLS file |
| write_VIEWS | Write a VIEWS control file for a CRAN-style package repository |

The terms of the vocabulary are stored in a DAG, which can be loaded as the serialized data object biocViewsVocab. For listing of available terms use function getSubTerms.

Further information is available in the following two vignettes:

| HOWTO-BCV | Basic package usage |
|-----------------|---|
| createReposHtml | Further information for repository admins |

Author(s)

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Maintainer: Biocore Team c/o BioC user list <bioconductor@stat.math.ethz.ch>

Examples

```
data(biocViewsVocab)
getSubTerms(biocViewsVocab, "Technology")
```

BiocView-class Class "BiocView"

Description

Representation of of Bioconductor "view".

Objects from the Class

Objects can be created by calls of the form new("BiocView", ...).

bioc Views Vocab

Slots

name: Object of class "character" giving the name of the view.

subViews: Object of class "character" giving the names of the subviews of this view.

parentViews: Object of class "character" giving the names of the views that are this view's parents.

Title: Object of class "character" giving longer description of view?

reposRoot: Object of class "character" URL for repository

homeUrl: Object of class "character" ?

htmlDir: Object of class "character" ?

packageList: Object of class "list" consisting of PackageDetail-class objects

Extends

Class "RepositoryDetail", directly. Class "Htmlized", directly.

Methods

```
coerce signature(from = "BiocView", to = "rdPackageTable"): ...
htmlDoc signature(object = "BiocView"): ...
htmlFilename signature(object = "BiocView"): ...
htmlValue signature(object = "BiocView"): ...
show signature(object = "BiocView"): ...
```

Author(s)

Seth Falcon

biocViewsVocab Bioconductor Task Views Vocabulary Data

Description

A graphNEL-class instance representing the Bioconductor Task Views as a directed graph.

Usage

```
data(biocViewsVocab)
```

Format

The format is: graphNEL instance

Details

The source for the vocabulary data is in the dot directory of the package in file biocViewsVocab.dot. This is transformed to GXL using the dot2gxl command line utility from the graphviz package. Then the fromGXL function from the graph package is used to convert to graphNEL-class.

Examples

data(biocViewsVocab) biocViewsVocab ## If you have Rgraphviz available, you can ## plot the vocabulary with plot(biocViewsVocab)

dump_concept produce character stream for use with Protege entities/class hierarchy ingestion based on biocViews subgraphs

Description

produce character stream for use with Protege entities/class hierarchy ingestion based on biocViews subgraphs

Usage

dump_concept(x, edob = edges(biocViewsVocab))

Arguments

| х | character(1) name of a node in the biocViewsVocab graph |
|------|---|
| edob | a list as produced by graph::edges |

Value

a character atom with x abutting left and edge targets tab-indented

Examples

cat(substring(dump_concept("ResearchField"), 1, 152), "\n")

extractManuals

Description

This function extracts Rd man pages and builds pdf reference manuals from the man subdirectory of R source packages archives (.tar.gz) found in a local package repository.

All Rd files found in man will be extracted and used during the pdf construction process. Only source package archives will be processed. The constructed pdf files will be extracted under destDir and will be found in PKGNAME/man/*.pdf.

Prior to extraction, all Rd and pdf files in destDir/PKGNAME/man will be removed.

Usage

extractManuals(reposRoot, srcContrib, destDir)

Arguments

| reposRoot | character vector giving the path to the root of the local CRAN-style package repository |
|------------|---|
| srcContrib | character vector giving the relative path from the reposRoot to the source pack- ages. In a standard CRAN-style repository, this will be src/contrib. |
| destDir | character vector specifying the directory in which the extracted files will be written. If missing, files will be written to <reposroot>/manuals.</reposroot> |

Author(s)

Patrick Aboyoun

```
extractNEWS
```

Extract NEWS files from source package tarballs

Description

Extracts NEWS files from source tarballs of packages.

Usage

```
extractNEWS(reposRoot, srcContrib, destDir)
```

Arguments

| reposRoot | Top level path for CRAN-style repos |
|------------|-------------------------------------|
| srcContrib | Location of source packages |
| destDir | where to extract |

extractTopLevelFiles Extract files from the top level of source package tarballs

Description

Extracts files from source tarballs of packages.

Usage

```
extractTopLevelFiles(reposRoot, srcContrib, destDir, fileName)
```

Arguments

| reposRoot | Top level path for CRAN-style repos |
|------------|-------------------------------------|
| srcContrib | Location of source packages |
| destDir | where to extract |
| fileName | name of file to extract |

extractVignettes Extract pdf vignettes from local package repository

Description

These functions extract pdf or HTML files from the inst/doc subdirectory of R source packages archives (.tar.gz) found in a local package repository.

All pdf files found in inst/doc will be extracted. With extractHTMLDocuments, all HTML files except index.html will be extracted. Only source package archives will be processed. The extracted pdf or HTML files will be extracted under destDir and will be found in PKGNAME/inst/doc/.

Prior to extraction, all pdf files in destDir/PKGNAME/inst/doc will be removed.

Usage

```
extractVignettes(reposRoot, srcContrib, destDir)
extractHTMLDocuments(reposRoot, srcContrib, destDir)
```

Arguments

| reposRoot | character vector giving the path to the root of the local CRAN-style package repository |
|------------|---|
| srcContrib | character vector giving the relative path from the reposRoot to the source pack- ages. In a standard CRAN-style repository, this will be src/contrib. |
| destDir | character vector specifying the directory in which the extracted files will be written. If missing, files will be written to <reposroot>/vignettes.</reposroot> |

genReposControlFiles

Author(s)

Seth Falcon

genReposControlFiles Generate CRAN-style repository control files

Description

This function generates control files for CRAN-style repositories. For each path specified in contribPaths a PACKAGES file is written. In addition, two top-level control files are created:

REPOSITORY contains information about the specified contrib paths.

VIEWS contains metadata for all packages in the repository including the paths to any extracted vignettes, if found. This file is useful for generating HTML views of the repository.

Usage

```
genReposControlFiles(reposRoot, contribPaths, manifestFile = NA,
  meatPath = NA)
```

Arguments

| reposRoot | character vector containing the path to the CRAN-style repository root directory. |
|--------------|--|
| contribPaths | A named character vector. Valid names are source, win.binary, mac.binary, mac.binary.mavericks, and mac.binary.el-capitan. Values indicate the paths to the package archives relative to the reposRoot. |
| manifestFile | character(1). File path location to Bioconductor formatted manifest file that lists all current packages. This file will be used in the write_VIEWS function to cross check successfully built packages with all expected packages. Packages that have not built will be given dummy entry for complete listing in bioc_VIEWS. If NA cross check is skipped and packages not built on any system will be missing from biocVIEWS |
| meatPath | character(1). File path location to the directory containing cloned repositories of Bioconductor packages. If manifestFile is used for cross checking and the meatPath is provided, entries from the DESCRIPTION file are manually entered into biocVIEWS information. If NA dummy values for minimal fields for landing page generation are included with ERROR. This attempts to fill in as much information as possible for packages that have failed to build. |

Author(s)

Seth Falcon

See Also

write_PACKAGES, extractVignettes, write_REPOSITORY, write_VIEWS

getBiocSubViews

Description

This function returns a list of BiocView-class objects corresponding to the subgraph of the views DAG induced by topTerm. In short, this does the same thing as getBiocViews, but limits the vocabulary to topTerm and all of its decendents.

Usage

```
getBiocSubViews(reposUrl, vocab, topTerm, local = FALSE, htmlDir = "")
```

Arguments

| reposUrl | URL for a CRAN-style repository that hosts a VIEWS file at the top-level. |
|----------|---|
| vocab | A graph-class object representing the ontologyof views. This graph should be a directed acyclic graph (DAG). |
| topTerm | A string giving the name of the subview DAG. This view and all of its decendents will be included in the result. |
| local | logical indicating whether to assume a local package repository. The default is FALSE in which case absolute links to package detail pages are created. |
| htmlDir | if the local argument is TRUE, this will be used as the relative path for package HTML files. |

Details

The root of the vocabulary DAG is implicitly included in the view creation process order to build views with a link back to the top. It is removed from the return list.

This function is tailored to generation of Bioconductor Task Views. With the current vocabulary, it probably only makes sense to call it with topView set to one of "Software", "AnnotationData", or "ExperimentData". This is a hack to allow the biocViews code to manage HTML views across more than one repository.

Value

A list of BiocView-class objects. The names of the list give the name of the corresponding view.

Author(s)

Seth Falcon

See Also

write_VIEWS, writeBiocViews

getBiocViews

Examples

```
data(biocViewsVocab)
reposPath <- system.file("doc", package="biocViews")
reposUrl <- paste("file://", reposPath, sep="")
biocViews <- getBiocSubViews(reposUrl, biocViewsVocab, "Software")
print(biocViews[1:2])</pre>
```

```
getBiocViews
```

Build a list of BiocView objects from a package repository

Description

Given the URL to a CRAN-style package repository containing a VIEWS file at the top-level and a graph-class object representing a DAG of views, this function returns a list of BiocView-class objects.

Usage

```
getBiocViews(reposUrl, vocab, defaultView, local = FALSE, htmlDir = "")
```

Arguments

| reposUrl | URL for a CRAN-style repository that hosts a VIEWS file at the top-level. |
|-------------|---|
| vocab | A graph-class object representing the ontology of views. This graph should be a directed acyclic graph (DAG). |
| defaultView | A string giving the term to use for packages that do not list a term of their own via the biocViews field in the 'DESCRIPTION' file. |
| local | logical indicating whether to assume a local package repository. The default is FALSE in which case absolute links to package detail pages are created. |
| htmlDir | if the local argument is TRUE, this will be used as the relative path for package HTML files. |

Value

A list of BiocView-class objects. The names of the list give the name of the corresponding view.

Author(s)

Seth Falcon

See Also

write_VIEWS, writeBiocViews

Examples

```
data(biocViewsVocab)
reposPath <- system.file("doc", package="biocViews")
reposUrl <- paste("file://", reposPath, sep="")
biocViews <- getBiocViews(reposUrl, biocViewsVocab, "NoViewProvided")
print(biocViews[1:2])</pre>
```

getCurrentbiocViews Get a list of biocViews for each branch

Description

This function looks returns a list containing all the biocViews that are present on the Bioconductor website.

Usage

```
getCurrentbiocViews()
```

Details

It parses the dot file present inside the biocViews package.

Value

It returns a named list with 3 components.

| Software | biocViews from the software branch |
|----------------|--|
| ExperimentData | biocViews from the ExperimentData branch |
| AnnotationData | biocViews from the AnnotationData branch |

Author(s)

Sonali Arora

Examples

```
ans <- getCurrentbiocViews()
## only the first 6 from each branch are shown here.
lapply(ans, head)</pre>
```

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getPackageNEWS

Description

These functions visit two Bioconductor releases, identifying packages that are present in the 'current' repository and have NEWS since the base version of the same package in the 'previous' release. All NEWS is reported for packages only in the current repository.

Usage

Arguments

| prevRepos | character(1) Bioconductor version from which NEWS starts. |
|--------------|---|
| currRepos | character(1) Bioconductor version for current packages. |
| геро | character(1) Which repository to get NEWS for. bioc is software packages, data/experiment is for data experiment packages and workflows for workflow packages |
| srcdir | Path to local checkout of package repositories, if NULL will try and use files on main builders |
| dbs | A list of news_db elements, as returned by getPackageNEWS. |
| destfile | character(1) file path to the location where NEWS will be printed. |
| overwrite | logical(1) indicating whether destfile can be over-written, if it exists. |
| width | numeric(1) number of characters news items are to be wrapped to, excluding indent. |
| output | character(1) output to text or markdown format. |
| relativeLink | Should links to packages be relative links on bioconductor.org website or include full url 'https//bioconductor.org'. default: FALSE is full url. |
| | additional arguments, unused. |

Value

A list of news_db files, as returned by utils::news, for each package for which relevant NEWS is available.

Author(s)

Martin Morgan mtmorgan@fhcrc.org and Lori Shepherd

getPackageTitles

Description

These functions visit two Bioconductor releases branches, identifying differnt packages that are present in the 'current' repository from the 'previous' release. Utilizes the devel branch of bioconductor to retrieve description.

Usage

Arguments

| prevBranch | character(1) Bioconductor branch to compare to |
|--------------|---|
| currBranch | character(1) Bioconductor branch for current packages. |
| manifest | character(1) Which repository of pakcages to compare. software.txt is software packages, data-experiment.txt is for data experiment packages and work-flows.txt for workflow packages |
| status | get new or removed package list comparing currBranch to prevBranch |
| pkgs | character() A list of packages to retrieve DESCRIPTION |
| outfile | character(1) file path to the location where DESCRIPTIONS will be printed. |
| output | character(1) output to text or markdown format. |
| relativeLink | Should links to packages be relative links on bioconductor.org website or include full url 'https//bioconductor.org'. default: FALSE is full url. |

Value

A list of package titles.

Author(s)

Martin Morgan mtmorgan@fhcrc.org and Lori Shepherd

getPacksAndViews

Examples

```
## Not run:
# At release time get a list of new or removed or deprecated packages
# get new packages in release 3.7 that are not in 3.6
newSoft = getPackageTitles()
# get removed packages from 3.6
rmSoft = getPackageTitles(currBranch="RELEASE_3_7", status="removed")
# get depreacted package for 3.7
deprecatedSoft = setdiff(getPackageTitles(status="removed"), rmSoft)
# repeated above for data-experiment packages
newData = getPackageTitles(manifest="data-experiment.txt")
rmData = getPackageTitles(currBranch="RELEASE_3_7", manifest="data-experiment.txt", status="removed"), rmData)
deprecatedData = setdiff(getPackageTitles(manifest="data-experiment.txt", status="removed"), rmData)
```

End(Not run)

getPacksAndViews Parse VIEWS file for views and packages

Description

Given a repository URL, download and parse the VIEWS file.

Usage

```
getPacksAndViews(reposURL, vocab, defaultView, local=FALSE)
```

Arguments

| reposURL | character vector giving the URL of a CRAN-style repository containing a VIEWS file at the top-level. |
|-------------|--|
| vocab | A graph-class object representing the ontologyof views. This graph should be a directed acyclic graph (DAG). |
| defaultView | A string giving the term to use for packages that do not list a term of their own via the biocViews field in the 'DESCRIPTION' file. |
| local | logical indicating whether certain links should be absolute (using reposURL) or relative. |

Value

A list with named elements:

views: Vector of view memberships. Names are package names.

pkgList: A list of PackageDetail-class objects.

Author(s)

Seth Falcon

getSubTerms

Retrieve a term and its children from a vocab DAG

Description

Given a Directed Acyclic Graph (DAG) represented as a graphNEL instance, return a character vector consisting of the specified term and all of its descendants. That is, give the list of terms for which a path exists starting at term.

Usage

getSubTerms(dag, term)

Arguments

| dag | A graphNEL representing a DAG |
|------|--|
| term | A string giving a term in the vocabulary (a node in dag) |

Value

A character vector of term names.

Author(s)

S. Falcon

Examples

```
data(biocViewsVocab)
getSubTerms(biocViewsVocab, "Software")
```

guessPackageType

Guess Package Type (Software, ExperimentData, AnnotationData) using existing biocViews.

Description

biocViews are "keywords" which are used to describe a given package. They are broadly divided into three categories, representing the type of packages present in the Bioconductor Project - Software, Annotation Data and Experiment Data. biocViews are supposed to come from only one of the three fields, but this function will check the list of biocViews and guess the package type based on how many biocViews came from each field.

Usage

guessPackageType(biocViews)

Arguments

biocViews A character vector containing a list of biocViews.

Value

A character(1) of package type: either "Software", "ExperperimentData", or "AnnotationData".

Author(s)

Lori Shepherd

Examples

```
guessPackageType(c("clustering", "classification"))
guessPackageType(c("Organism", "Homo Sapien"))
```

htmlDoc

Create a complete HTML document representation of an object

Description

This generic function should return an XMLNode instance representing the specified object in HTML as a complete HTML document.

Usage

htmlDoc(object, ...)

Arguments

| object | An object |
|--------|---------------------|
| | Not currently used. |

Value

An instance of XMLNode from the XML package.

Author(s)

Seth Falcon

See Also

htmlValue, htmlFilename

htmlFilename

Return a filename for an object's HTML representation

Description

This function returns a string containing an appropriate filename for storing the object's HTML representation.

Usage

htmlFilename(object, ...)

Arguments

| object | An object. |
|--------|--------------------|
| | Not currently used |

Value

A character vector of length one containing the filename.

Author(s)

Seth Falcon

See Also

htmlValue, htmlDoc

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Htmlized-class Class "Htmlized"

Description

A virtual class for HTML serialization method dispatch.

Objects from the Class

A virtual Class: No objects may be created from it.

Methods

htmlDoc signature(object = "Htmlized"): Return the html-ized representation of object as a complete HTML document.

Author(s)

Seth Falcon

htmlValue

HTML Representation of an Object

Description

This generic function should return an XMLNode instance representing the specified object in HTML

Usage

```
htmlValue(object)
```

Arguments

object An object

Value

An instance of XMLNode from the XML package.

Author(s)

Seth Falcon

See Also

htmlDoc, htmlFilename

PackageDetail-class Class "PackageDetail"

Description

Representation of R package metadata. Most slots correspond to fields in a package's DESCRIP-TION file.

Objects from the Class

Objects can be created by calls of the form new("PackageDetail", ...).

Slots

Package: Object of class "character" see DESCRIPTION Version: Object of class "character" see DESCRIPTION Title: Object of class "character" see DESCRIPTION Description: Object of class "character" see DESCRIPTION Author: Object of class "character" see DESCRIPTION Maintainer: Object of class "character" see DESCRIPTION Depends: Object of class "character" see DESCRIPTION Imports: Object of class "character" see DESCRIPTION Suggests: Object of class "character" see DESCRIPTION SystemRequirements: Object of class "character" see DESCRIPTION License: Object of class "character" see DESCRIPTION URL: Object of class "character" see DESCRIPTION biocViews: Object of class "character" see DESCRIPTION vignettes: Object of class "character" giving paths to vignette pdf files in the repository vignetteScripts: Object of class "character" giving paths to vignette Stangled R files in the repository vignetteTitles: Object of class "character" giving the titles of the vignette files in the repository source.ver: Object of class "character" version string for the source package win.binary.ver: Object of class "character" version string for the 32-bit Windows binary package mac.binary: Object of class "character" version string for the macOS High Sierra binary package mac.binary.mavericks.ver: Object of class "character" version string for the OS X Mavericks binary package mac.binary.el-capitan.ver: Object of class "character" version string for the OS X El Capitan binary package

- downloadStatsUrl: Object of class "character" An optional URL for the download history statistics.
- manuals: Object of class "character" giving paths to reference manual pdf files in the repository
- dependsOnMe: Object of class "character" giving packages found in the repository that depend on this package
- importsMe: Object of class "character" giving packages found in the repository that imports this package
- suggestsMe: Object of class "character" giving packages found in the repository that suggest this package
- functionIndex: Object of class "character" Not used. Intended to hold function index data.
- reposFullUrl: Object of class "character" The URL for the full URL of the root of the repository.

reposRoot: Object of class "character" The URL for the root of the repository.

viewRoot: Object of class "character" The URL for the view of the repository.

devHistoryUrl: Object of class "character" The URL for the development changelog.

Extends

Class "Htmlized", directly.

Methods

- **htmlDoc** signature(object = "PackageDetail"): Return an XMLNode instance containg a complete HTML document representation of the package.
- **htmlFilename** signature(object = "PackageDetail"): Return a filename appropriate for the HTML document representation.
- htmlValue signature(object = "PackageDetail"): Return XMLNode instance containing an HTML
 representation of the package.

Details

pdAuthorMaintainerInfo-class pdVignetteInfo-class pdDownloadInfo-class pdDetailsInfo-class pdDescriptionInfo-class pdVigsAndDownloads-class

Dummy classes for HTML generation. Each dummy class is a simple extension (it does not add any slots). The purpose of each dummy class is to allow for method dispatch to generate HTML via the htmlValue method.

You can convert convert a PackageDetail instance to one of the dummy classes like this: descInfo <- as(pdObj, "pdDescriptionInfo")

Author(s)

Seth Falcon

Examples

```
pd <- new("PackageDetail",</pre>
          Package="MyFancyPackage",
          Version="1.2.3",
          Title="A Fancy Package",
          Description="This package does fancy things",
          Author="A. Coder",
          Maintainer="A. Coder <acoder@foo.bar.net>",
          Depends="methods",
          Imports="ASimplePackage",
          Suggests="MyDataPackage",
          biocViews="Infrastructure",
      vignettes="vignettes/MyFancyPackage/inst/doc/MFP1.pdf,\nvignettes/MyFancyPackage/inst/doc/MFP2.pdf",
      vignetteScripts="vignettes/MyFancyPackage/inst/doc/MFP1.R\nvignettes/MyFancyPackage/inst/doc/MFP2.R",
          vignetteTitles="MFP1 Document,\nMFP2 Document",
          source.ver="src/contrib/MyFancyPackage_1.2.3.tar.gz",
          win.binary.ver="bin/windows/contrib/4.0/MyFancyPackage_1.2.2.zip",
          mac.binary.ver="bin/macosx/contrib/4.0/MyFancyPackage_1.2.3.tgz",
          dependsOnMe=c("PackageThatExposesMe"),
          importsMe=c("AnEvenFancierPackage","AMuchFancierPackage"),
          suggestsMe="PackageThatUsesMeInVignette",
          reposRoot="http://foo.bar.org")
html <- htmlValue(pd)</pre>
pd
```

recommendBiocViews Recommend biocViews for an existing Package.

Description

Packages being added to the Bioconductor Project require biocViews in their DESCRIPTION file.(Note that the field name "biocViews" is case-sensitive and must begin with a lower-case 'b'.)biocViews are "keywords" which are used to describe a given package. They are broadly divided into three categories, representing the type of packages present in the Bioconductor Project - Software, Annotation Data and Experiment Data.

Usage

recommendBiocViews(pkgdir, branch)

Arguments

| pkgdir | The path of the package Directory. | |
|--------|--|-----|
| branch | The branch which your package will belong to. It can be either 'Software | e', |
| | 'AnnotationData' or 'ExperimentData'. | |

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recommendPackages

Details

This function parses the package directory provided by the user to recommend biocViews to the user. The output is a suggested list - the user of this function is expected to go through this list and find which biocViews best describe his or her package. It uses the following strategies.

- It parses the "Description", "Title", "Package" of the DESCRIPTION page to find biocViews.
- It looks up the biocViews of the packages in the "Depends" field of the given package to recommend biocViews
- It parses the text from the man pages and the vignettes to suggest biocViews.

Please note the following:

- Do not make up your own biocViews.
- Double check the spelling and case of the biocViews added.
- Please add biocViews only from the appropriate branch. eg: Software packages should have only Software biocViews.

Value

A list is returned with 3 characters - current, recommended and remove.

- "current" contains the biocViews from the package's DESCRIPTION file.
- "recommended" are the recommended biocViews This is a suggested list which the user can add in addition to "current" biocViews the user is expected to go through this list and find which biocViews best describe their package.
- "remove" are those biocViews which are inconsistent with the Bioconductor biocViews. (Hint check for spelling, cases and plural)

Author(s)

Sonali Arora.

recommendPackages Recommend Packages using existing biocViews.

Description

biocViews are "keywords" which are used to describe a given package. They are broadly divided into three categories, representing the type of packages present in the Bioconductor Project - Software, Annotation Data and Experiment Data. One can find packages which are tagged with biocViews using this function.

Usage

recommendPackages(biocViews, use.release=TRUE, intersect.views=TRUE)

Arguments

| biocViews | A character vector containing a list of biocViews. Currently only biocViews from the software branch are supported. | |
|-----------------|--|--|
| use.release | A logical character indicating if you want packages recommended from the re- lease branch of Biocondutor. | |
| intersect.views | | |
| | A logical character indicating if you want packages which are tagged with all the input biocViews or packages tagged with any one or more of the biocViews. | |

Value

A character vector containing a list of packages. If multiple biocViews are given as input, the result returns packages tagged with all or atleast one of the input biocViews.

Author(s)

Sonali Arora.

Examples

recommendPackages(c("clustering", "classification"))

RepositoryDetail-class

Class "RepositoryDetail"

Description

Representation of R package repository index

Objects from the Class

Objects can be created by calls of the form new("RepositoryDetail", ...).

Slots

Title: Object of class "character" giving the title for the repository.
reposRoot: Object of class "character" giving the root URL of the repository
homeUrl: Object of class "character" ?
htmlDir: Object of class "character" ?
packageList: Object of class "list" consisting of objects of class PackageDetail-class

Extends

Class "Htmlized", directly.

validate_bioc_views

Methods

htmlDoc signature(object = "RepositoryDetail"): ... htmlFilename signature(object = "RepositoryDetail"): ... htmlValue signature(object = "RepositoryDetail"): ...

Author(s)

Seth Falcon

validate_bioc_views Validate a package's biocViews.

Description

Ensures that a package has biocViews and that they are valid. Function is designed to be called from the unit tests of another package.

Usage

```
validate_bioc_views(pkg)
```

Arguments

pkg character(1) Name of package to validate.

Value

invisible(NULL) if tests pass.

Author(s)

Dan Tenenbaum dtenenba@fhcrc.org

Examples

validate_bioc_views("biocViews")

writeBiocViews

Description

This function serializes a list of BiocView-class objects to a series of HTML files.

Usage

```
writeBiocViews(bvList, dir, backgroundColor="transparent")
```

Arguments

| bvList | A list of BiocView-class objects |
|-----------------|---|
| dir | A character vector giving the directory where the HTML files will be written. |
| backgroundColor | |
| | A character vector giving the background color for the body in the CSS file. |

Author(s)

Seth Falcon

See Also

getBiocViews, genReposControlFiles, write_VIEWS

| writeHtmlDoc | Write an XML DOM containing HTML to a file | |
|--------------|--|--|
|--------------|--|--|

Description

Given a DOM tree from the XML package and a filename, write the DOM to disk creating an HTML file.

Usage

```
writeHtmlDoc(html, file)
```

Arguments

| html | A DOM object from the XML package |
|------|-----------------------------------|
| file | A string giving the filename |

Author(s)

S. Falcon

writePackageDetailHtml

Write HTML files for packages in a CRAN-style repository

Description

This function creates package "homepages" that describe the package and provide links to download package artifacts in the repository.

Usage

```
writePackageDetailHtml(pkgList, htmlDir = "html", backgroundColor="transparent")
```

Arguments

| pkgList | A list of PackageDescription objects. |
|-----------------|---|
| htmlDir | The files will be written to this directory. |
| backgroundColor | |
| | A character vector giving the background color for the body in the CSS file |

A character vector giving the background color for the body in the CSS file.

Author(s)

Seth Falcon

See Also

writeRepositoryHtml

writeRepositoryHtml Write package descriptions and a repository index as HTML

Description

This function generates an HTML file for each package in a repository and generates an index.html file that provides an alphabetized listing of the packages.

Usage

```
writeRepositoryHtml(reposRoot, title, reposUrl = "..", viewUrl = "../..",
    reposFullUrl=reposUrl, downloadStatsUrl="",
    devHistoryUrl="", link.rel = TRUE,
    backgroundColor="transparent")
```

Arguments

| reposRoot | string specifying the path to the root of the CRAN-style package repository. |
|------------------|--|
| title | string giving the title for the repository |
| reposUrl | string giving the prefix for URL in links generated on the package description pages. The default is "" which works well if the package description HTML files are written to an html subdirectory under the root of the repository. |
| viewUrl | string giving the prefix for the URL in links to the view pages. The biocViews terms will be linked to views summary pages with this prefix. |
| reposFullUrl | string giving the full prefix for URL in links generated on the package description pages. The default is reposUr1. |
| downloadStatsUrl | |
| | string giving the prefix for the URL in links to the download history statistics pages. |
| devHistoryUrl | string giving the prefix for the URL in links to the development changelog. |
| link.rel | logical indicating whether the index page should generate relative URL links. The default is TRUE. If you are generating HTML for a remote repository, you will want to set this to FALSE. |
| backgroundColor | |
| | A character vector giving the background color for the body in the CSS file. |

Author(s)

Seth Falcon

writeRFilesFromVignettes

Write R files from vignettes

Description

Ensures that .R files from vignette code chunks are written out.

Usage

Arguments

| reposRoot | Root directory of a CRAN-style repository |
|--------------|---|
| reposUrl | URL of repository |
| viewUrl | url of VIEWS file |
| reposFullUrl | Full URL of VIEWS file |

writeTopLevelView

downloadStatsUrl URL to download stats page devHistoryUrl Dev history URL

writeTopLevelView Write the view for the root of a vocabulary to disk

Description

Given a directory and a vocabulary represented as a graphNEL containing a DAG of terms, write the top-level term to disk as HTML.

This assumes your vocabulary has a single term with no parents.

Usage

writeTopLevelView(dir, vocab)

Arguments

| dir | A string giving a directory in which to write the HTML file |
|-------|---|
| vocab | A graphNEL instance giving the DAG of terms. It should have a root node. That |
| | is, there should be exactly one node with no incoming edges. |

Author(s)

S. Falcon

write_REPOSITORY Write a REPOSITORY control file for a CRAN-style package repository

Description

This function writes a REPOSITORY file at the top-level of a CRAN-style repository. This file is DCF formatted and describes the location of packages available in the repository. Here is an example for a repository containing source packages, and Windows and Mac binary packages:

```
source: src/contrib
win.binary: bin/windows/contrib/4.0
mac.binary: bin/macosx/contrib/4.0
provides: source, win.binary, mac.binary
```

Usage

write_REPOSITORY(reposRootPath, contribPaths)

Arguments

| reposRootPath | character vector containing the path to the CRAN-style repos | sitory root directory. |
|---------------|--|------------------------|
| contribPaths | A named character vector. Valid names are source, win.bi | nary, mac.binary, |
| | <pre>mac.binary.mavericks, and mac.binary.el-capitan.</pre> | Values indicate the |
| | paths to the package archives relative to the reposRoot. | |

Author(s)

Seth Falcon

See Also

write_PACKAGES, extractVignettes, genReposControlFiles, write_VIEWS

| write_SYMBOLS | Write a SYMBOLS file | |
|---------------|----------------------|--|
|---------------|----------------------|--|

Description

Writes a DCF formatted file, SYMBOLS, containing the symbols exported by each package in a directory containg R package source directories.

Usage

```
write_SYMBOLS(dir, verbose = FALSE, source.dirs=FALSE)
```

Arguments

| dir | The root of a CRAN-style package repository containing source packages. When source.dirs is TRUE, dir should be a directory containing R package source directories |
|-------------|--|
| verbose | Logical. When TRUE, progress is printed to the standard output. |
| source.dirs | Logical. When TRUE, interpret dir as a directory containing source package directories. When FALSE, the default, dir is assumed to be the root of a CRAN-style package repository and the function will operate on the source package tarballs in dir/src/contrib. |

Value

Returns NULL. Called for the side-effect of creating a SYMBOLS file in dir.

Author(s)

S. Falcon

See Also

write_PACKAGES write_VIEWS

Description

This function writes a VIEWS file to the top-level of a CRAN-style package repository. The VIEWS file is in DCF format and describes all packages found in the repository.

The VIEWS file contains the complete DESCRIPTION file for each source package in the repository. In addition, metadata for available binary packages and vignettes is centralized here.

Usage

```
write_VIEWS(reposRootPath, fields = NULL,
            verbose = FALSE, vignette.dir = "vignettes",
            manifestFile = NA, meatPath = NA)
```

Arguments

| reposRootPath | character vector containing the path to the CRAN-style repository root directory. |
|---------------|---|
| fields | Any additional fields to include. You shouldn't need this, but if you have added fields to the DESCRIPTION files of the packages in the repository, you may want it. |
| verbose | logical, if TRUE, print progress messages. |
| vignette.dir | character specifying where to look for vignettes. |
| manifestFile | character(1). File path location to Bioconductor formatted manifest file that lists all current packages. This file will be used in the write_VIEWS function to cross check successfully built packages with all expected packages. Packages that have not built will be given dummy entry for complete listing in bioc_VIEWS. If NA cross check is skipped and packages not built on any system will be missing from biocVIEWS |
| meatPath | character(1). File path location to the directory containing cloned repositories of Bioconductor packages. If manifestFile is used for cross checking and the meatPath is provided, entries from the DESCRIPTION file are manually en- tered into biocVIEWS information. If NA dummy values for minimal fields for landing page generation are included with ERROR. This attempts to fill in as much information as possible for packages that have failed to build. |

Warning

This function uses a private function from the tools package: tools:::.build_repository_package_db.

Author(s)

Seth Falcon

See Also

write_PACKAGES, extractVignettes, genReposControlFiles, write_REPOSITORY

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