

Package ‘AnnotationHubData’

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Description These recipes convert a wide variety and a growing number of public bioinformatic data sets into easily-used standard Bioconductor data structures.

License Artistic-2.0

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makeUCSCChain.R makeUCSC2Bit.R makeUCSCTracks.R
trackWithAuxiliaryTableToGRangesRecipe.R
UCSCTrackUpdateChecker.R makeEnsemblTwoBit.R

NeedsCompilation no

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AnnotationHubRecipes-package

Transform public data resources into Bioconductor Data Structures
 ~~ package title ~~

Description

These recipes convert a wide variety and a growing number of public bioinformatic data sets into easily-used standard Bioconductor data structures.

Details

Package: AnnotationHubRecipes
 Type: Package
 Version: 1.0
 Date: 2012-11-30
 License: Artistic-2.0

This package provides a set of methods which convert bioinformatic data resources into standard Bioconductor data types. For example, a UCSC genome browser track, expressed as a BED file, is converted into a GRanges object. Not every valuable data resource can be transformed quite so easily; some require more elaborate transformation, and hence a more specialized recipe. Every effort is made to limit the number of recipes required. One strategy that helps with the principle of "zero curation": unless absolutely required, the "cooked" version of the data resource produced by a recipe is a simple and unembellished reflection of the original data in its downloaded form.

Author(s)

Dan Tenenbaum, Paul Shannon

AnnotationHubMetadata-class

Class "AnnotationHubMetadata" and methods

Description

AnnotationHubMetadata is used to represent record(s) in the server data base.

Usage

```
AnnotationHubMetadata(AnnotationHubRoot, SourceUrl, SourceType,
  SourceVersion, SourceLastModifiedDate, SourceMd5 =
  NA_character_, SourceSize, DataProvider, Title,
  Description, Species, TaxonomyId, Genome, Tags,
  Recipe, RDataClass, RDataDateAdded, RDataPath,
  Maintainer, ..., BiocVersion = biocVersion(),
  Coordinate_1_based = TRUE, Notes = NA_character_,
  DispatchClass, Location_Prefix =
  "http://s3.amazonaws.com/annotationhub/")

jsonPath(x)
toJson(x)
constructAnnotationHubMetadataFromSourceFilePath(ahroot, originalFile)
constructMetadataFromJsonPath(ahroot, jsonpath)
constructSeqInfo(species, genome)

metadata(x, ...)
hubError(x)
inputFiles(object, ...)
outputFile(object)
ahmToJson(ahm)
deleteResources(id)
getImportPreparerClasses()
makeAnnotationHubResource(objName, makeAnnotationHubMetadataFunction,
  ..., where)
```

Arguments

AnnotationHubRoot	character(1) Absolute path to directory structure containing resources to be added to AnnotationHub. Internal use only.
SourceUrl	character() URL of original resource(s).
SourceType	character() Form of original data, e.g., BED, FASTA, etc.
SourceVersion	character(1) Version of original file.
SourceLastModifiedDate	POSIXct() The date when the source was last modified.

SourceMd5	character()	md5 hash of original file.
SourceSize	numeric(1)	Size of original file in bytes.
DataProvider	character(1)	Provider of original data, e.g., NCBI, UniProt etc.
Title	character(1)	Title for the resource with version or genome build as appropriate.
Description	character(1)	Description of the resource. May include details such as data type, format, study origin, sequencing technology, treated vs control, number of samples etc.
Species	character(1)	Species name.
TaxonomyId	character(1)	NCBI code
Genome	character(1)	Name of genome build.
Tags	character()	Free-form tags that serve as search terms.
Recipe	character(1)	Name of recipe function. Only applicable to recipes created by the Bioconductor core team and included in AnnotationHubData base code.
RDataClass	character()	Class of derived R object, e.g., GRanges. Length must match the length of RDataPath.
RDataDateAdded	POSIXct()	Date resource was added to AnnotationHub; used to determine snapshots.
RDataPath	character()	File path to serialized form. Length must match the length of RDataClass. Internal use only.
Maintainer	character(1)	Maintainer name and email address, e.g., 'YourName YourName@email.com'
BiocVersion	A character(1)	Bioconductor version. The resource will be available in Bioconductor \geq to this version. Default value is the current version, specified with biocVersion().
Coordinate_1_based	logical(1)	Do coordinates start with 1 or 0?
DispatchClass	character(1)	String used in AnnotationHub:::get1() method dispatch when loading the resource from AnnotationHub. This value is often the same as the RDataClass. Dispatch classes are defined on the .get1() methods in AnnotationHub. If one exists for your data type, simply use the name of the data type, e.g., if the data are saved as ExpressionSet objects, DispatchClass should be "ExpressionSet"; when the resource is downloaded from AnnotationHub with '[' the "Resources" extension is (automatically) added to make the new class "ExpressionSetResource". If the data are saved as .rda files DispatchClass should be "Rda" and when downloaded in AnnotationHub the class name becomes "RdaResource" and so on. If there is not a .get1() method in AnnotationHub for your object type, contact maintainer@bioconductor.org for assistance.
Location_Prefix	character(1)	Base path where the resource is coming from, e.g., a web site or Amazon S3.
Notes	character()	Notes about the resource.
ahm		An instance of class AnnotationHubMetadata.
x		An instance of class AnnotationHubMetadata.

jsonpath	character(1) Full path to a JSON representation of AnnotationHubMetadata-class.
ahroot	A character(1) AnnotationHubRoot to be added to the path / location of the returned instance.
originalFile	A character(1) Original file name.
object	An AnnotationHubRecipe instance.
species	character(1) The organism, e.g., "Homo sapiens".
genome	character(1) The genome build, e.g., "hg19".
id	An id whose DB record is to be fully deleted.
objName	character(1) The name of the PreparerClass used for dispatch.
makeAnnotationHubMetadataFunction	function Function (name) that makes AnnotationHubMetadata objects from the resource(s).
where	Environment where function definition is defined. Default value is sufficient.
...	Additional arguments passed to methods.

Value

AnnotationHubMetadata returns an instance of the class.

jsonPath returns a character(1) representation of the full path to the location of the json file associated with this record.

toJson returns the JSON representation of the record.

fromJson returns an instance of the class, as parsed from the JSON file.

Objects from the Class

Objects can be created by calls to the constructor, AnnotationHubMetadata().

Author(s)

Dan Tenenbaum and Marc Carlson

Examples

```
getClass("AnnotationHubMetadata")
```

flog

flog

Description

Write logging message to console and a file.

Usage

```
flog(level, ...)
```

Arguments

level A character(1) string object.
... Further arguments.

Details

Writes the message to the console and to a file.

Value

None.

Author(s)

Dan Tenenbaum

ImportPreparer-class *Class ImportPreparer and generic newResources*

Description

The ImportPreparer and derived classes are used for dispatch during data discovery (see [newResources](#)). There is one ImportPreparer class for each data source for [AnnotationHubMetadata](#).

`newResources` is a generic function; with methods implemented for each ImportPreparer.

Author(s)

Martin Morgan

See Also

[AnnotationHubMetadata](#).

Examples

```
getImportPreparerClasses()
```

`makeAnnotationHubMetadata`*Make AnnotationHubMetadata objects from csv file of metadata*

Description

Make AnnotationHubMetadata objects from .csv files located in the "inst/extdata/" package directory of an AnnotationHub package.

Usage

```
makeAnnotationHubMetadata(pathToPackage, fileName=character())
```

Arguments

`pathToPackage` Full path to data package including the package name; no trailing slash

`fileName` Name of metadata file(s) with csv extension. If none are provided, all files with .csv extension in "inst/extdata" will be processed.

Details

- `makeAnnotationHubMetadata`: Reads the resource metadata from .csv files into a [AnnotationHubMetadata](#) object. The [AnnotationHubMetadata](#) is inserted in the AnnotationHub database. Intended for internal use or package authors checking the validity of package metadata.

Value

A named list the length of `fileName`. Each element is a list of of AnnotationHubMetadata objects created from the .csv file.

See Also

- [updateResources](#)
- [readMetadataFromCsv](#)
- [AnnotationHubMetadata](#) class

Examples

```
## Not run:  
makeAnnotationHubMetadata("path/to/mypackage")  
  
## End(Not run)
```

makeEnsemblFasta	<i>Functions to convert Ensembl FASTA files to FaFile and TwoBitFile for inclusion in AnnotationHub.</i>
------------------	--

Description

Transform an Ensembl FASTA file to a Bioconductor FaFile or ToBitFile.

Usage

```
makeEnsemblFastaToAHM(currentMetadata, baseUrl = "ftp://ftp.ensembl.org/pub/",
                      baseDir = "fasta/", release,
                      justRunUnitTest = FALSE, BiocVersion = biocVersion())

makeEnsemblTwoBitToAHM(currentMetadata, baseUrl = "ftp://ftp.ensembl.org/pub/",
                       baseDir = "fasta/", release,
                       justRunUnitTest = FALSE, BiocVersion = biocVersion())

ensemblFastaToFaFile(ahm)

ensemblFastaToTwoBitFile(ahm)
```

Arguments

currentMetadata	Currently not used. Intended to be a list of metadata to filter, i.e., records that do not need to be processed again. Need to remove or fix.
baseUrl	ftp file location.
baseDir	ftp file directory.
release	Integer version number, e.g., "84".
justRunUnitTest	A logical. When TRUE, a small number of records (usually 5) are processed instead of all.
BiocVersion	A character(1) Bioconductor version. The resource will be available in Bioconductor >= to this version. Default value is the current version, specified with biocVersion().
ahm	List of AnnotationHubMetadata instances.

Details

makeEnsemblFastaToAHM and makeEnsemblTwoBitToAHM process metadata into a list of AnnotationHubMetadata objects.

ensemblFastaToFaFile unzips a .gz files, creates an index and writes out .rz and .rz.fai files to disk. ensemblFastaToTwoBit converts a fasta file to twobit format and writes the .2bit file out to disk.

Value

makeEnsemblFastaToAHM and makeEnsemblTwoBitToAHM return a list of AnnotationHubMetadata objects.

ensemblFastaToFaFile write out .rz and .rz.fai files to disk. ensemblFastaToTwoBit writes out a .2bit file to disk.

Author(s)

Bioconductor Core Team

See Also

- [updateResources](#)
- [AnnotationHubMetadata](#)

Examples

```
## updateResources() generates metadata, process records and
## pushes files to AWS S3 buckets. See ?updateResources for details.

## 'release' is passed to makeEnsemblFastaToFaFile.
## Not run:
meta <- updateResources("/local/path",
                        BiocVersion = c("3.2", "3.3"),
                        preparerClasses = "EnsemblFastaImportPreparer",
                        metadataOnly = TRUE, insert = FALSE,
                        justRunUnitTest = FALSE, release = "83")

## End(Not run)
```

makeGencodeFasta

Recipe to add Gencode FASTA resources to AnnotationHub

Description

Create metadata and process raw Gencode FASTA files for inclusion in AnnotationHub

Usage

```
makeGencodeFastaToAHM(currentMetadata,
                      baseUrl="ftp://ftp.sanger.ac.uk/pub/gencode/",
                      species=c("Human", "Mouse"), release,
                      justRunUnitTest=FALSE,
                      BiocVersion=biocVersion())

gencodeFastaToFaFile(ahm)
```

Arguments

currentMetadata	Currently not used. Intended to be a list of metadata to filter, i.e., records that do not need to be processed again. Need to remove or fix.
baseUrl	ftp file location.
species	A character(1) of the species. Currently "Human" and "Mouse" are supported.
release	A character string of the release number.
justRunUnitTest	A logical. When TRUE, a small number of records (usually 5) are processed instead of all.
BiocVersion	A character vector of Bioconductor versions the resources should be available for.
ahm	List of AnnotationHubMetadata instances.

Details

- Documentation: <http://www.gencodegenes.org/releases/>
- File download location: <ftp://ftp.sanger.ac.uk/pub/gencode/>. Gencode_human and Gencode_mouse are used.
- Files downloaded: Code is currently specific for human and mouse. Files chosen for download are described in AnnotationHubData:::gencodeDescription().

Value

makeGencodeFastaAHM returns a list of AnnotationHubMetadata instances. gencodeFastaToFastaFile returns nothing.

Author(s)

Bioconductor Core Team.

See Also

- [updateResources](#)
- [AnnotationHubMetadata](#)

Examples

```
## updateResources() generates metadata, process records and
## pushes files to AWS S3 buckets.

## To run the GencodeFasta recipe specify
## 'preparerClasses = GencodeFastaImportPreparer'. The 'species' and 'release'
## arguments are passed to makeGencodeFastaAHM().
## Not run:
meta <- updateResources("/local/path",
                        BiocVersion = c("3.2", "3.3"),
                        preparerClasses = "GencodeFastaImportPreparer",
                        metadataOnly = TRUE, insert = FALSE,
                        justRunUnitTest = FALSE)
```

```
## End(Not run)
```

```
makeStandardOrgDbs    Functions to add OrgDb and TxDb sqlite files to AnnotationHub
```

Description

Add OrgDb and TxDb sqlite files to AnnotationHub

Usage

```
makeStandardOrgDbsToAHM(currentMetadata, justRunUnitTest = FALSE,
                        BiocVersion = biocVersion(),
                        downloadOrgDbs = TRUE)

makeStandardTxDbsToAHM(currentMetadata, justRunUnitTest = FALSE,
                       BiocVersion = biocVersion(), TxDbs)

makeNCBIToOrgDbsToAHM(currentMetadata, justRunUnitTest = FALSE,
                      BiocVersion = biocVersion(),
                      baseUrl = "ftp://ftp.ncbi.nlm.nih.gov/gene/DATA/")
```

Arguments

currentMetadata	Historically was intended to be a list of metadata to filter, i.e., records that do not need to be processed again. In some recipes this is used as a way to pass additional arguments. Need to remove or make consistent.
baseUrl	A character(). The file location.
justRunUnitTest	A logical. When TRUE, a small number of records (usually <= 5) are processed instead of all.
BiocVersion	A character(1). The resource will be available for Bioconductor versions greater than and equal to this version. Default is BiocInstaller::biocVersion().
TxDbs	Character vector of the TxDb names; generally includes TxDbs that were new or updated for the current release.
downloadOrgDbs	A logical. Indicates if all OrgDb packages in the Bioconductor repo should be downloaded and installed. This should be TRUE the first time the recipe is run and can be FALSE for subsequent runs when testing.

Details

makeStandardOrgDbsToAHM and makeStandardTxDbsToAHM extracts the sqlite files from the existing OrgDb and TxDb packages in the Bioconductor repositories and generate associated metadata.

makeNCBIToOrgDbsToAHM creates sqlite files and metadata for 1000 organisms with the makeOrgPackageFromNCBI function. These organisms are less 'main stream' than those hosted in the Bioconductor repository (makeStandardOrgDbsToAHM) and the databases are less comprehensive because data only come from one source, NCBI.

Value

List of AnnotationHubMetadata objects.

Author(s)

Bioconductor Core Team

See Also

- [updateResources](#)
- [AnnotationHubMetadata](#)

Examples

```
## Not run:
## In Bioconductor 3.5, one new TxDb was added and 4 active
## tracks were updated. This piece of code shows how to add these 5
## packages to AnnotationHub.

## Step I: generate metadata
##
## Generate the metadata with the low-level helper for inspection.
TxDb1 <- c("TxDb.Ggallus.UCSC.galGal5.refGene",
           "TxDb.Celegans.UCSC.ce11.refGene",
           "TxDb.Rnorvegicus.UCSC.rn5.refGene",
           "TxDb.Dmelanogaster.UCSC.dm6.ensGene",
           "TxDb.Rnorvegicus.UCSC.rn6.refGene")
meta <- makeStandardTxDbToAHM(currentMetadata=list(AnnotationHubRoot=getwd()),
                             justRunUnitTest=FALSE,
                             TxDb1 = TxDb1)

## Once the low-level helper runs with no errors, try generating the
## metadata with the high-level wrapper updateResources(). Setting
## metadataOnly=TRUE will generate metadata only and not push resources
## to S3. insert=FALSE prevents the metadata from being inserted in the
## database.
##
## The metadata generated by updateResources() will be the same as that
## generated by makeStandardTxDbToAHM(). Both should be a list the same
## length as the number of TxDb1 specified.
meta <- updateResources(getwd(),
                       preparerClasses="TxDbFromPkgsImportPreparer",
                       metadataOnly=TRUE, insert = FALSE,
                       justRunUnitTest=FALSE, TxDb1 = TxDb1)

INFO [2017-04-11 09:12:09] Preparer Class: TxDbFromPkgsImportPreparer
complete!
> length(meta)
[1] 5

## Step II: push resources to S3
##
## If the metadata looks correct we are ready to push resources to S3.
## Set metadataOnly=FALSE but keep insert=FALSE.

meta <- updateResources(getwd(),
```

```

        BiocVersion="3.5",
        preparerClasses="TxDbFromPkgsImportPreparer",
        metadataOnly=FALSE, insert = FALSE,
        justRunUnitTest=FALSE, TxDBs = TxDBs)

## Step III: insert metadata in AnnotationHub production database
##
## Inserting the metadata in the database is usually done as a separate step
## and with the help of the AnnotationHub docker.
## Set metadataOnly=TRUE and insert=TRUE.
meta <- updateResources(getwd(),
                        BiocVersion="3.5",
                        preparerClasses="TxDbFromPkgsImportPreparer",
                        metadataOnly=FALSE, insert = FALSE,
                        justRunUnitTest=FALSE, TxDBs = TxDBs)

## End(Not run)

```

readMetadataFromCsv *Reads csv files of resource metadata into a data.frame*

Description

Reads metadata files located in the "inst/extdata" package directory into a data.frame.

Usage

```
readMetadataFromCsv(pathToPackage, fileName=character())
```

Arguments

`pathToPackage` Full path to data package including package name; no trailing slash.

`fileName` Name of single metadata file located in "inst/extdata". If none is provided the function looks for a file named "metadata.csv".

Details

- readMetadataFromCsv:

Reads a .csv file of metadata located in "inst/extdata". readMetadataFromCsv performs checks for required columns and data types and can be used by package authors to validate their metadata before submitting the package for review. The function is used internally by makeAnnotationHubMetadata.

The rows of the .csv file(s) represent individual Hub resources (i.e., data objects) and the columns are the metadata fields. All fields should be a single character string of length 1.

Required Fields in metadata file:

- Title: character(1). Name of the resource. This can be the exact file name (if self-describing) or a more complete description.
- Description: character(1). Brief description of the resource, similar to the 'Description' field in a package DESCRIPTION file.

- `BiocVersion`: character(1). The first Bioconductor version the resource was made available for. Unless removed from the hub, the resource will be available for all versions greater than or equal to this field.
- `Genome`: character(1). Genome.
- `SourceType`: character(1). Format of original data, e.g., FASTA, BAM, BigWig, etc.
- `SourceUrl`: character(1). Optional location of original data files. Multiple urls should be provided as a comma separated string.
- `SourceVersion`: character(1). Version of original data.
- `Species`: character(1). Species.
- `TaxonomyId`: character(1). Taxonomy ID.
- `Coordinate_1_based`: logical. TRUE if data are 1-based.
- `DataProvider`: character(1). Name of company or institution that supplied the original (raw) data.
- `Maintainer`: character(1). Maintainer name and email in the following format: Maintainer Name <username@address>.
- `RDataClass`: character(1). R / Bioconductor class the data are stored in, e.g., GRanges, SummarizedExperiment, ExpressionSet etc.
- `DispatchClass`: character(1). Determines how data are loaded into R. The value for this field should be 'Rda' if the data were serialized with `save()` and 'Rds' if serialized with `saveRDS`. The filename should have the appropriate 'rda' or 'rds' extension. A number of dispatch classes are pre-defined in `AnnotationHub/R/AnnotationHubResource-class.R` with the suffix 'Resource'. For example, if you have sqlite files, the `AnnotationHubResource-class.R` defines `SQLiteFileResource` so the `DispatchClass` would be `SQLiteFile`. Contact `maintainer@bioconductor.org` if you are not sure which class to use.
- `ResourceName`: character(1). Exact file name.
- `Location_Prefix`: character(1). Do not include this field if data are stored in AWS S3; it will be generated automatically. If data will be accessed from a location other than AWS S3 this field should be the base url. When `Location_Prefix` is provided you also need to specify `RDataPath`.
- `RDataPath`: character(1). Do not include this field if data are stored in AWS S3; it will be generated automatically. If data will be accessed from a location other than AWS S3 this field should be the remainder of the path to the resource. The `Location_Prefix` will be prepended to `RDataPath` for the full path to the resource.
- `Tags`: character() vector. 'Tags' are search terms used to define a subset of resources in a Hub object, e.g. in a call to query. For ExperimentHub resources, 'Tags' are automatically generated from the 'biocViews' in the DESCRIPTION. 'Tags' values supplied by the user are not be entered in the database and are not part of the formal metadata. This 'controlled vocabulary' approach was taken to limit the search terms to a well defined set and may change in the future. 'Tags' for AnnotationHub resources are a free-form field of search terms defined by the user. The package name is added as one of the 'Tags' before the metadata are finalized. Multiple 'Tags' are specified as a colon separated string, e.g., tags for two resources would look like this:

```
Tags=c("tag1:tag2:tag3", "tag1:tag3")
```

NOTE: The metadata file can have additional columns beyond the 'Required Fields' listed above. These values are not added to the Hub database but they can be used in package functions to provide an additional level of metadata on the resources.

Value

A data.frame with one row per resource and columns for the Required Fields described above. Additional auto-generated columns, e.g., RDataDateAdded and PreparerClass may also be present and are used by internal functions when generating the final metadata.

See Also

- [makeAnnotationHubMetadata](#)

Examples

```
## Each row of the metadata file represents a resource added to one of
## the 'Hubs'. This example creates a metadata.csv file for a single resource.
## In the case of multiple resources, the arguments below would be character
## vectors that produced multiple rows in the data.frame.
```

```
meta <- data.frame(
  Title = "RNA-Sequencing dataset from study XYZ",
  Description = paste0("RNA-seq data from study XYZ containing 10 normal ",
    "and 10 tumor samples represented as a",
    "SummarizedExperiment"),
  BiocVersion = "3.4",
  Genome = "GRCh38",
  SourceType = "BAM",
  SourceUrl = "http://www.path/to/original/data/file",
  SourceVersion = "Jan 01 2016",
  Species = "Homo sapiens",
  TaxonomyId = 9606,
  Coordinate_1_based = TRUE,
  DataProvider = "GEO",
  Maintainer = "Your Name <youremail@provider.com>",
  RDataClass = "SummarizedExperiment",
  DispatchClass = "Rda",
  ResourceName = "FileName.rda"
)

## Not run:
## Write the data out and put in the inst/extdata directory.
write.csv(meta, file="metadata.csv", row.names=FALSE)

## Test the validity of metadata.csv with readMetadataCsv():
readMetadataFromCsv("path/to/mypackage")

## End(Not run)
```

 updateResources

updateResources

Description

Add new resources to AnnotationHub

Usage

```
updateResources(AnnotationHubRoot, BiocVersion = biocVersion(),
               preparerClasses = getImportPreparerClasses(),
               metadataOnly = TRUE, insert = FALSE,
               justRunUnitTest = FALSE, ...)

pushResources(allAhms, uploadToS3 = TRUE, download = TRUE)

pushMetadata(allAhms, url)
```

Arguments

AnnotationHubRoot	Local path where files will be downloaded.
BiocVersion	A character(1) Bioconductor version. The resource will be available in Bioconductor \geq to this version. Default value is the current version, specified with <code>biocVersion()</code> .
preparerClasses	One of the <code>ImportPreparer</code> subclasses defined in <code>getImportPreparer()</code> . This class is used for dispatch during data discovery.
metadataOnly	A logical to specify the processing of metadata only or both metadata and data files. When <code>FALSE</code> , metadata are generated and data files are downloaded, processed and pushed to their final location in S3 buckets. <code>metadata = TRUE</code> produces only metadata and is useful for testing.
insert	NOTE: This option is for inserting metadata records in the production data base (done by Bioconductor core team member) and is for internal use only. A logical to control if metadata are inserted in the AnnotationHub db. By default this option is <code>FALSE</code> which is a useful state in which to test a new recipe and confirm the metadata fields are correct. When <code>insert = TRUE</code> , the "AH_SERVER_POST_URL" global option must be set to the http location of the AnnotationHubServer in the global environment or <code>.Rprofile</code> . Additionally, AWS command line tools must be installed on the local machine to push files to S3 buckets. See https://aws.amazon.com/cli/ for installation instructions.
justRunUnitTest	A logical. When <code>TRUE</code> , a small number of records (usually 5) are processed instead of all.
allAhms	List of <code>AnnotationHubMetadata</code> objects.
url	URL of AnnotationHub database where metadata will be inserted.
uploadToS3	A logical indicating whether resources should be uploaded to the AWS S3 bucket.
download	A logical indicating whether resources should be downloaded from resource url.
...	Arguments passed to other methods such as <code>regex</code> , <code>baseUrl</code> , <code>baseDir</code> .

Details

- `updateResources`:

updateResources is responsible for creating metadata records and downloading, processing and pushing data files to their final resting place. The

- preparerClasses argument is used in method dispatch to determine which recipe is used. By manipulating the metadataOnly, insert and justRunUnitTest arguments one can flexibly test the metadata for a small number of records with or without downloading and processing the data files.
- global options:
Several recipes look at the "AnnotationHub_Use_Disk" option to determine if the file is written to S3. This is legacy behavior and not clearly documented. If the recipe being run respects this option it must be set before running updateResources,
When insert = TRUE the "AH_SERVER_POST_URL" option must be set to the https location of the AnnotationHub db.

Value

A list of AnnotationHubMetadata objects.

Author(s)

Martin Morgan, Marc Carlson

See Also

- [AnnotationHubMetadata](#)

Examples

```
## Not run:

## -----
## Inspect metadata:
## -----
## A useful first step in testing a new recipe is to generate and
## inspect a small number of metadata records. The combination of
## 'metadataOnly=TRUE', 'insert=FALSE' and 'justRunUnitTest=TRUE'
## generates metadata for the first 5 records and does not download or
## process any data.

meta <- updateResources("/local/path",
                        BiocVersion = "3.3",
                        preparerClasses = "EnsemblFastaImportPreparer",
                        metadataOnly = TRUE, insert = FALSE,
                        justRunUnitTest = TRUE,
                        release = "84")

INFO [2015-11-12 07:58:05] Preparer Class: EnsemblFastaImportPreparer
Ailuropoda_melanoleuca.ailMel1.cdna.all.fa.gz
Ailuropoda_melanoleuca.ailMel1.dna_rm.toplevel.fa.gz
Ailuropoda_melanoleuca.ailMel1.dna_sm.toplevel.fa.gz
Ailuropoda_melanoleuca.ailMel1.dna.toplevel.fa.gz
Ailuropoda_melanoleuca.ailMel1.ncrna.fa.gz

## The return value is a list of metadata for the first 5 records:
```

```

> names(meta)
[1] "FASTA cDNA sequence for Ailuropoda melanoleuca"
[2] "FASTA DNA sequence for Ailuropoda melanoleuca"
[3] "FASTA DNA sequence for Ailuropoda melanoleuca"
[4] "FASTA DNA sequence for Ailuropoda melanoleuca"
[5] "FASTA ncRNA sequence for Ailuropoda melanoleuca"

## Each record is of class AnnotationHubMetadata:

> class(meta[[1]])
[1] "AnnotationHubMetadata"
attr(,"package")
[1] "AnnotationHubData"

## -----
## Insert metadata in the db and process/push data files:
## -----
## This next code chunk creates the metadata and downloads and processes
## the data (metadataOnly=FALSE). If all files are successfully pushed to
## to their final resting place, metadata records are inserted in the
## AnnotationHub db (insert=TRUE). Metadata insertion is done by a
## Bioconductor team member; contact maintainer@bioconductor.org for help.

meta <- updateResources("local/path",
                        BiocVersion = "3.5",
                        preparerClasses = "EnsemblFastaImportPreparer",
                        metadataOnly = FALSE, insert = TRUE,
                        justRunUnitTest = FALSE,
                        regex = ".*release-81")

## -----
## Recovery helpers:
## -----

## pushResources() and pushMetadata() are both called from updateResources()
## but can be used solo for testing or completing a run that
## terminated unexpectedly.

## Download, process and push to S3 the last 2 files in 'meta':
sub <- meta[length(meta) - 1:length(meta)]
pushResources(sub)

## Insert metadata in the AnotationHub db for the last 2 files in 'meta':

pushMetadata(sub, url = getOption("AH_SERVER_POST_URL"))

## End(Not run)

```

upload_to_S3

Upload a file to Amazon S3

Description

This function is for uploading a file resource to the S3 cloud.

Usage

```
upload_to_S3(file, remotename, bucket, profile, acl="public-read")
```

Arguments

file	The file to upload.
remotename	The name this file should have in S3, including any "keys" that are part of the name. This should not start with a slash (if it does, the leading slash will be removed), but can contain forward slashes.
bucket	Name of the S3 bucket to copy to.
profile	Corresponds to a profile set in the config file for the AWS CLI (see the documentation). If this argument is omitted, the default profile is used.
acl	Should be one of private, public-read, or public-read-write.

Details

Uses the [AWS Command Line Interface](#) to copy a file to Amazon S3. Assumes the CLI is properly configured and that the aws program is in your PATH. The CLI should be configured with the credentials of a user who has permission to upload to the appropriate bucket. It's recommended to use [IAM](#) to set up users with limited permissions.

There is an `RAmazonS3` package but it seems to have issues uploading files to S3.

Value

TRUE on success. If the command fails, the function will exit with an error.

Author(s)

Dan Tenenbaum

Examples

```
## Not run:
upload_to_S3("myfile.txt", "foo/bar/baz/yourfile.txt")
# If this is successful, the file should be accessible at
# http://s3.amazonaws.com/annotationhub/foo/bar/baz/yourfile.txt

## End(Not run)
```

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