Package 'ArrayExpress'

December 9, 2025

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Title Access the ArrayExpress Collection at EMBL-EBI Biostudies and build Bioconductor data structures: ExpressionSet, AffyBatch, NChannelSet
Version 1.71.0
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Depends R (>= 2.9.0), Biobase (>= 2.4.0)
Imports oligo, limma, httr, utils, jsonlite, rlang, tools, methods
Description Access the ArrayExpress Collection at EMBL- EBI Biostudies and build Bioconductor data structures: ExpressionSet, AffyBatch, NChannelSet
License Artistic-2.0
biocViews Microarray, DataImport, OneChannel, TwoChannel
NeedsCompilation no
Suggests affy
git_url https://git.bioconductor.org/packages/ArrayExpress
git_branch devel
git_last_commit 4a25159
git_last_commit_date 2025-10-29
Repository Bioconductor 3.23
Date/Publication 2025-12-09
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ae2bioc

Convert MAGE-TAB files from raw data into a Bioconductor object

Description

ae2bioc converts local MAGE-TAB files into a AffyBatch, an ExpressionSet or a NChannelSet.

Usage

```
ae2bioc(mageFiles, dataCols = NULL, drop = TRUE)
```

Arguments

mageFiles A list as given from getAE function. Containing the following elements:

rawFiles all the expression files to use to create the object. The content of the raw.zip MAGE-TAB file.

sdrf the name of the sdrf file from MAGE-TAB. **idf** the name of the idf file from MAGE-TAB.

adf the name of the adf file from MAGE-TAB.

dataCols by default, the columns are automatically selected according to the scanner type.

path is the name of the directory containing these files.

If the scanner is unknown or if the user wants to use different columns than the default, the argument 'dataCols' can be set. For two colour arrays it must be a list with the fields 'R', 'G', 'Rb' and 'Gb' giving the column names to be used for red and green foreground and background. For one colour arrays, it must be a character string with the column name to be used. These column names must

correspond to existing column names of the expression files.

drop if TRUE and only one platform in series, the platform name will be dropped.

Value

An object of class AffyBatch, ExpressionSet or NChannelSet with the raw expression values in the 'assayData' of the object, the information contained in the sdrf file in the 'phenoData', the adf file content in the 'featureData' and the idf file content in the 'experimentData'.

If several array designs are used in the dataset, the output is a list with an object for each array design.

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See Also

ArrayExpress, queryAE, getAE

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Examples

An example can be found in the help of the getAE function.

ArrayExpress R objects from ArrayExpress collection

Description

ArrayExpress produces an AffyBatch, an ExpressionSet or a NChannelSet from a raw dataset from the ArrayExpress collection of the Biostudies database. ArrayExpress needs an Internet connection.

Usage

ArrayExpress(accession, path = tempdir(), save = FALSE, dataCols = NULL, drop = TRUE)

Arguments

accession an ArrayExpress experiment identifier. path the name of the directory in which the files downloaded on the ArrayExpress repository will be extracted. The default is the current directory. if TRUE, the files downloaded from the database will not be deleted from path save after executing the function. dataCols by default, for the raw data, the columns are automatically selected according to the scanner type. If the scanner is unknown or if the user wants to use different columns than the default, the argument 'dataCols' can be set. For two colour arrays it must be a list with the fields 'R', 'G', 'Rb' and 'Gb' giving the column names to be used for red and green foreground and background. For one colour arrays, it must be a character string with the column name to be used. These column names must correspond to existing column names of the expression files. if TRUE and only one platform in series, the platform name will be dropped. drop

Value

The output is an object of class AffyBatch or ExpressionSet or NChannelSet with the raw expression values in the assayData of the object, the information contained in the .sdrf file in the phenoData, the adf file in the featureData and the idf file content in the experimentData.

If several array designs are used in the data set, the output is a list with an object for each array design.

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See Also

```
queryAE, getAE, ae2bioc, getcolproc, procset
```

Examples

```
ETABM25.affybatch = ArrayExpress("E-TABM-25")
print(ETABM25.affybatch)
sampleNames(ETABM25.affybatch)
colnames(pData(ETABM25.affybatch))
```

extract.zip

Unzip archives in a specified directory

Description

extract.zip extracts the files from a .zip archive in a specific directory.

Usage

```
extract.zip(file, extractpath = dirname(file)[1])
```

Arguments

file A file name.

extractpath A path to define where the files are to be extracted.

Value

Success is indicated by returning the directory in which the files have been extracted. If it fails, it returns an empty character string.

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getAE	Download MAGE-TAB files from the ArrayExpress collection in a
	specified directory

Description

getAE downloads and extracts the MAGE-TAB files from an ArrayExpress dataset.

Usage

```
getAE(accession, path = getwd(), type = "full", extract = TRUE, sourcedir = path, overwrite = FALSE)
```

Arguments

accession	is an ArrayExpress experiment identifier.
path	is the name of the directory in which the files downloaded on the ArrayExpress repository will be extracted.
type	can be 'raw' to download and extract only the raw data, 'processed' to download and extract only the processed data or 'full' to have both raw and processed data.
extract	if FALSE, the files are not extracted from the zip archive.
sourcedir	when local = TRUE, files will be read from this directory.
overwrite	if TRUE, overwrite files if they already exist in path, default FALSE.

Value

A list with the names of the files that have been downloaded and extracted.

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```

See Also

```
ArrayExpress, ae2bioc, getcolproc, procset
```

Examples

```
mexp21 = getAE("E-MEXP-21", type = "full")

## Build a an ExpressionSet from the raw data
MEXP21raw = ae2bioc(mageFiles = mexp21)

## Build a an ExpressionSet from the processed data
cnames = getcolproc(mexp21)
MEXP21proc = procset(mexp21, cnames[2])
```

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getcolproc

Return the possible column names from processed MAGE-TAB files

Description

getcolproc extracts the column names from processed MAGE-TAB and return them. The output is needed to call the function procset.

Usage

```
getcolproc(files)
```

Arguments

files

A list as given from getAE function. Containing the following elements:

procfile procfile is the name of the processed MAGE-TAB file to be read.

path is the name of the directory where to find this file.

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See Also

ArrayExpress, queryAE, getAE, procset

getcolraw

Return the possible column names from raw MAGE-TAB files

Description

getcolraw extracts the column names from raw MAGE-TAB and return them. The output can be use to set the argument 'rawcol' of the function magetab2bioc.

Usage

```
getcolraw(rawfiles)
```

Arguments

rawfiles

rawfiles are the name of the raw MAGE-TAB files to be read.

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See Also

ArrayExpress, queryAE, getAE

procset

Convert processed MAGE-TAB files into a Bioconductor object

Description

procset converts local MAGE-TAB files into an ExpressionSet.

Usage

```
procset(files, procol)
```

Arguments

files is the list with the names of the processed, the sdrf, the adf and the idf files and

the path of the data as given by getAE.

procol the name of the column to be extracted from the file. Obtained using getcolproc.

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See Also

```
queryAE, getAE, getcolproc
```

Examples

An example can be found in the help of the getAE function.

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queryAE

Query of the ArrayExpress collection in Biostudies

Description

queryAE queries the ArrayExpress collection with keywords and give a dataframe with ArrayExpress identifiers and related information, as an output.

Usage

```
queryAE(keywords = NULL, species = NULL)
```

Arguments

keywords the keyword(s) of interest. To use several words, they must be separated by a

"+" as shown in the examples.

species the specie(s) of interest.

Value

A dataframe with all the ArrayExpress dataset identifiers which correspond to the query in the first column. The following columns contain information about these datasets, such as the number of files, the release date on the database, the title, the author and content.

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See Also

```
ArrayExpress, getAE
```

Examples

```
## To retrieve all the identifiers of pneumonia data sets
pneumo = queryAE(keywords = "pneumonia")

## To retrieve all the identifiers of pneumonia data sets studied in human
pneumoHS = queryAE(keywords = "pneumonia", species = "homo+sapiens")
```

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