# Package 'makecdfenv'

October 16, 2019
<b>Version</b> 1.60.0
<b>Date</b> 2006-03-06
Title CDF Environment Maker
Author Rafael A. Irizarry <rafa@jhu.edu>, Laurent Gautier</rafa@jhu.edu>
<laurent@cbs.dtu.dk>, Wolfgang Huber</laurent@cbs.dtu.dk>
<w.huber@dkfz-heidelberg.de>, Ben Bolstad</w.huber@dkfz-heidelberg.de>
Maintainer James W. MacDonald < jmacdon@u.washington.edu>
<b>Depends</b> R ( $\geq$ 2.6.0), affyio
Imports Biobase, affy, methods, stats, utils, zlibbioc
<b>Description</b> This package has two functions. One reads a Affymetrix chip description file (CDF) and creates a hash table environment containing the location/probe set membership mapping. The other creates a package that automatically loads that environment.
License GPL (>= 2)
LazyLoad yes
biocViews OneChannel, DataImport, Preprocessing
git_url https://git.bioconductor.org/packages/makecdfenv
git_branch RELEASE_3_9
git_last_commit 900ece3
git_last_commit_date 2019-05-02
Date/Publication 2019-10-15
R topics documented:
Cdf-class getInfoInFile make.cdf.env make.cdf.package pmormm read.cdffile
Index

2 Cdf-class

Cdf-class

Class Cdf

#### **Description**

A class describing the content a Chip Description File.

#### **Details**

The class Cdf was designed to store the content of Affymetrix's Chip Definition Files (CDF). This early class is no longer widely used in the package affy. Environments (called cdfenvs) are preferred to have an efficient mapping between probe set identifiers and indexes (needed to access/subset particular probe intensities). Unless one needs to access every information contained in a CDF file, the cdfenvs will be preferred.

The following comments are only relevant to someone with interest in what is in a CDF file. The name associated to each probe is not unique, it corresponds to the gene name. It is very common to have a name repeated 40 times (20 perfect matches and 20 mismatches). Probes can be grouped by pairs: a perfect match (PM) probe has its mismatch (MM) counterpart. The two probes in a pair differ by one base (usually located in the middle of the sequence). The information relative to this particular base are stored in pbase and tbase. At a given position (x,y), having pbase[x,y] == tbase[x,y] means having a MM while having pbase[x,y] != tbase[x,y] means having a perfect mismatch if and only if one of the bases is A while the other is TRUE, or one is G and the other is C. The function pmormm returns TRUE for PMs and FALSE for MMs.

To know more about the file structure of a CDF file, one has to refer to the parsing code (mostly in C).

# **Creating Objects**

```
new('Cdf',
cdfName = ..., # Object of class character
name = ..., # Object of class matrix
name.levels = ..., # Object of class character
pbase = ..., # Object of class matrix
pbase.levels = ..., # Object of class character
tbase = ..., # Object of class matrix
tbase.levels = ..., # Object of class character
atom = ..., # Object of class matrix
```

#### **Slots**

```
cdfName: the CDF name tag. Used to link with the right CEL files
name: Object of class "matrix" of 'factors' for the gene names corresponding to the probes.
name.levels: Object of class "character" containing the levels corresponding to name.
pbase: Object of class "matrix" of pbase levels.
pbase.levels: Object of class "character" containing the levels corresponding to pbase.
tbase: Object of class "matrix" of tbase levels.
tbase.levels: Object of class "character" containing the levels corresponding to tbase.
atom: Object of class "matrix" of atom or probe numbers.
```

getInfoInFile 3

#### Methods

atom (Cdf): An accesor function for the ato, slot.

atom<- (Cdf): A replacement function for atom slot.

name.levels (Cdf): An accesor function for the name.levels slot.

**name.levels<-** (Cdf): A replacement function for name.levels slot.

pbase (Cdf): An accesor function for the pbase slot.

**pbase<-** (Cdf): A replacement function for pbase slot.

pbase.levels (Cdf): An accesor function for the pbase.levels slot.

pbase.levels<- (Cdf): A replacement function for pbase.levels slot.

show (Cdf): renders information about the Cdf object in a concise way on stdout.

tbase (Cdf): An accesor function for the tbase slot.

**tbase<-** (Cdf): A replacement function for tbase slot.

tbase.levels (Cdf): An accesor function for the tbase.levels slot.

tbase.levels<- (Cdf): A replacement function for tbase.levels slot.

#### Author(s)

L. Gautier < laurent@cbs.dtu.dk>

#### See Also

```
read.cdffile, make.cdf.env, make.cdf.package
```

getInfoInFile

get information from Affymetrix data files

# **Description**

Get specific information stored in Affymetrix data files (CEL or CDF).

# Usage

```
getInfoInFile(filename, type, unit, property, compress = NULL)
```

# **Arguments**

filename a file name

type "CEL" or "CDF" are the only known types.

unit the 'unit' to find the information

property the 'property' of interest

compress a boolean

4 make.cdf.env

#### **Details**

The data files seem to have a structure of 'units'. A 'unit' start with something between square brackets (ex: "[UNIT1]"). The parameter unit let one specify in which unit the information of interest is located. The 'properties' are constituted of an identifier, the sign '=' and the value for the property (ex: "numCells=12000").

If you are planning to use this function, I assume you know the what is in CEL and CDF files in details and that you know what are doing.

## Value

a character with the value of the 'property'.

#### Note

The code goes through the file until the rigth 'unit' is found. Then it looks for the right 'property' further down (eventually going through the next units if the property is not found).

#### See Also

```
read.celfile, read.cdffile, whatcdf
```

make.cdf.env

CDF Environment Maker

# Description

Reads an Affymetrix chip description file (CDF) and creates an environment used as a hash table for the probe set mapping to location.

# Usage

# Arguments

filename Character. Filename of the CDF file - **without** the path prefix!

cdf.path Character. Path to the CDF file.

compress Logical. If TRUE, CDF file is compressed.

return.env.only

Logical. If TRUE (the default), then the function returns an environment. Otherwise, a list with two elements, the first being the environment, and the second being a data structure that contains additional information needed for the pack-

age builder (see details, and vignette).

verbose Logical. If TRUE, messages are shown.

make.cdf.package 5

#### **Details**

Normally, this function should not be called directly. The preferred way to handle CDF information is to use make.cdf.package to build a package, and to install it into R. The CDF information can then either be invoked automatically by the package affy, or can be loaded manually by calling, for example, library(hgu133a).

Some R installations (typically on Windows) do not offer all the tools that are necessary for package building. In such situations, this function may be called directly. Please see the vignette for details - type:

```
openVignette("makecdfenv")
```

**Return values:** env is an environment, used as a hash table. For every probe set name we have a matrix with 2 columns. The first column contains the PM locations and the second column the MM locations. For PM only chips the MM column will have NAs.

syms is a list that contains chip-specific (i.e., CDF-file specific) information that can be used in the construction of the help files for the CDF package.

#### Value

Depending on the argument return.env.only, either the environment, or a list with two elements, env and syms. See details.

### Author(s)

Rafael A. Irizarry, Wolfgang Huber

#### See Also

```
make.cdf.package
```

#### **Examples**

make.cdf.package

CDF Environment Package Maker

# Description

This function reads an Affymetrix chip description file (CDF) and creates an R package that when loaded has the CDF environment available for use.

6 make.cdf.package

#### Usage

```
make.cdf.package(filename,
 packagename = NULL,
 cdf.path
              = getwd(),
 package.path = getwd(),
  compress
             = FALSE,
              = "The Bioconductor Project",
  author
 maintainer = "Biocore Package Maintainer <maintainer@bioconductor.org>",
             = packageDescription("makecdfenv", fields ="Version"),
 version
              = NULL,
  species
              = FALSE,
 unlink
              = TRUE)
 verbose
```

# Arguments

filename Character. Filename of the CDF file - without the path prefix!

packagename Character. Name wanted for the package.

cdf.path Character. Path to the CDF file.

package.path Character. Path where the package will be created.

compress Logical. If TRUE, CDF file is compressed.

author Character. What to put in the author field of the package.

maintainer Character. What to put in the maintainer field of the package.

version Character. What to put in the version field. Should be a of the form x.x.x.

species Character. Must be specified using the format e.g., Homo\\_sapiens

unlink Logical. If TRUE, and a package directory exists already in package.path, that

is overwritten.

verbose Logical. If TRUE messages are shown.

#### **Details**

The function is called for its side effect, creating a package. By default the package name will be the name of the CDF file made lower case and with special characters removed (i.e. only alphanumeric).

In general one would want to use the name given in by cleancdfname(abatch@cdfName) with abatch an AffyBatch object obtained, for example, using ReadAffy. This is the package name that the affy package looks for by default.

If the user has a CEL file, called filename, the recommended package name for the environment is cleancdfname(whatcdf(filename)). This usually coincides with the default.

Please see the vignette for more details.

# Value

If success, the function returns the name of the created package.

# Author(s)

Rafael A. Irizarry, Wolfgang Huber

pmormm 7

#### See Also

```
make.cdf.env
```

# **Examples**

pmormm

PM or MM

# **Description**

Determing if intensities on an array from the cdffile are PM, MM, or neither.

# Usage

```
pmormm(cdf)
```

# **Arguments**

cdf

A Cdf object

#### **Details**

The intensities on a chip fall in three categories: perfect match (PM), mismatch (MM) or unknown (No information about them is contained in the CDF file, yet they were found informative (details to come)). PM are coded as TRUE, MM as FALSE and unknown as NA.

# Value

```
A matrix (see section 'details').
```

# Author(s)

L. Gautier < laurent@cbs.dtu.dk>

8 read.cdffile

read.cdffile

Read a CDF file

# **Description**

Read the data contained in a CDF file

#### Usage

```
read.cdffile(file, compress=FALSE)
```

#### **Arguments**

file the name of the CDF file

compress whether the file is compressed or not

#### **Details**

This function is intended for use by make.cdf.env, which in turn is called by make.cdf.package. User may not have much benefit from calling this function directly.

In order to save memory, the name corresponding to each value in the CEL is a factor. As in R factor objects cannot be also of type matrix, the names corresponding to the indices were stored in a vector of type character called name.levels. The same thing was done with the pbase and cbase information (more for consistency than by conviction that some memory could be saved here).

# Value

Returns a Cdf-class object.

#### Note

A Cdf object is not a cdf environment, which will be needed for the computation of expression values from the probe intensities in a AffyBatch.

#### Author(s)

Laurent Gautier (laurent@cbs.dtu.dk)

# See Also

```
Cdf-class
```

# **Examples**

```
fn <- system.file("extdata", "Hu6800.CDF.gz", package="makecdfenv") mycdf <- read.cdffile(fn, compress=TRUE) mycdf \  \  \,
```

# **Index**

```
*Topic file
    read.cdffile, 8
*Topic manip
    getInfoInFile, 3
    make.cdf.env, 4
    make.cdf.package, 5
    pmormm, 7
*Topic methods
    Cdf-class, 2
AffyBatch, 6
atom (Cdf-class), 2
atom, Cdf-method (Cdf-class), 2
atom<- (Cdf-class), 2
atom<-,Cdf-method(Cdf-class), 2
Cdf-class, 2
getInfoInFile, 3
make.cdf.env, 3, 4, 7, 8
make.cdf.package, 3, 5, 5, 8
name.levels(Cdf-class), 2
name.levels, Cdf-method (Cdf-class), 2
name.levels<- (Cdf-class), 2
name.levels<-,Cdf-method(Cdf-class), 2</pre>
pbase (Cdf-class), 2
pbase,Cdf-method(Cdf-class), 2
pbase.levels (Cdf-class), 2
pbase.levels, Cdf-method (Cdf-class), 2
pbase.levels<- (Cdf-class), 2</pre>
pbase.levels<-,Cdf-method(Cdf-class), 2</pre>
pbase<- (Cdf-class), 2</pre>
pbase<-,Cdf-method(Cdf-class), 2</pre>
pmormm, 2, 7
read.cdffile, 3, 8
ReadAffy, 6
show, Cdf-method (Cdf-class), 2
tbase (Cdf-class), 2
tbase, Cdf-method (Cdf-class), 2
```

```
tbase.levels(Cdf-class), 2
tbase.levels,Cdf-method(Cdf-class), 2
tbase.levels<-(Cdf-class), 2
tbase.levels<-,Cdf-method(Cdf-class), 2
tbase<-(Cdf-class), 2
tbase<-,Cdf-method(Cdf-class), 2
```