

oligoClasses

March 18, 2010

DBPDInfo-class *Class "DBPDInfo"*

Description

A class for Platform Design Information objects, stored using a database approach

Objects from the Class

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

Slots

`getdb`: Object of class "function"

`tableInfo`: Object of class "data.frame"

`manufacturer`: Object of class "character"

`genomebuild`: Object of class "character"

`geometry`: Object of class "integer" with length 2 (rows x columns)

Methods

No methods defined with class "DBPDInfo" in the signature.

FeatureSet-class *"FeatureSet" and "FeatureSet" Extensions*

Description

Classes to store data from Expression/Exon/SNP/Tiling arrays at the feature level.

Objects from the Class

The FeatureSet class is VIRTUAL. Therefore users are not able to create instances of such class.

Objects for FeatureSet-like classes can be created by calls of the form: `new(CLASSNAME, assayData, manufacturer, platform, exprs, phenoData, featureData, experimentData, annotation, ...)`. But the preferred way is using parsers like [read.celfiles](#) and [read.xysfiles](#).

Slots

```
manufacturer: Object of class "character"  
assayData: Object of class "AssayData"  
phenoData: Object of class "AnnotatedDataFrame"  
featureData: Object of class "AnnotatedDataFrame"  
experimentData: Object of class "MIAME"  
annotation: Object of class "character"  
.__classVersion__: Object of class "Versions"
```

Methods

```
show signature(.Object = "FeatureSet"): show object contents
```

Author(s)

Benilton Carvalho

See Also

[eSet](#), [VersionedBiobase](#), [Versioned](#)

Examples

```
set.seed(1)  
tmp <- 2^matrix(rnorm(100), ncol=4)  
rownames(tmp) <- 1:25  
colnames(tmp) <- paste("sample", 1:4, sep="")  
efs <- new("ExpressionFeatureSet", exprs=tmp)
```

mmindex

Accessors for PM, MM or background probes indices.

Description

Extracts the indexes for PM, MM or background probes.

Usage

```
mmindex(object, ...)  
pmindex(object, ...)  
bgindex(object, ...)
```

Arguments

```
object      FeatureSet or DBPDInfo object  
...         Extra arguments, not yet implemented
```

Details

The indices are ordered by 'fid', i.e. they follow the order that the probes appear in the CEL/XYS files.

Value

A vector of integers representing the rows of the intensity matrix that correspond to PM, MM or background probes.

Examples

```
## How pm() works
## Not run:
x <- read.celfiles(list.celfiles())
pms0 <- pm(x)
pmi <- pmindex(x)
pms1 <- exprs(x)[pmi,]
identical(pms0, pms1)

## End(Not run)
```

 geometry

Array Geometry Information

Description

For a given array, `geometry` returns the physical geometry of it.

Usage

```
geometry(object)
```

Arguments

object PDInfo object

Examples

```
if (require(pd.mapping50k.xba240))
  geometry(pd.mapping50k.xba240)
```

 QuantificationSet-class

Class "QuantificationSet"

Description

A virtual class to store summarized measures.

Objects from the Class

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

Slots

assayData: Object of class "AssayData"
 phenoData: Object of class "AnnotatedDataFrame"
 featureData: Object of class "AnnotatedDataFrame"
 experimentData: Object of class "MIAME"
 annotation: Object of class "character"
 .__classVersion__: Object of class "Versions"

Extends

Class `eSet`, directly. Class `VersionedBiobase`, by class "eSet", distance 2. Class `Versioned`, by class "eSet", distance 3.

Methods

No methods defined with class "QuantificationSet" in the signature.

See Also

[SnpQSet](#), [SnpCnvQSet](#)

Examples

```
showClass("QuantificationSet")
```

 mmSequence

Probe Sequences

Description

Accessor to the (PM/MM/background) probe sequences.

Usage

```
mmSequence(object)
pmSequence(object, ...)
bgSequence(object, ...)
```

Arguments

object FeatureSet, AffySNPPDInfo or DBPDInfo object
 ... additional arguments

Value

A DNASringSet containing the PM/MM/background probe sequence associated to the array.

calls *Accessors for SnpCallSet*

Description

Accessors and replacement methods for genotype calls and associated confidences.

Usage

```
calls(object)
callsConfidence(object)
calls(object) <-value
callsConfidence(object) <-value
```

Arguments

object	SnpCallSet object.
value	matrix

Value

Matrix with R rows (SNP's) and C columns (Samples) with genotype calls (1 - AA; 2 - AB; 3 - BB) or confidences.

See Also

[crlmm](#)

SnpCallSetPlus-class
Class "SnpCallSetPlus"

Description

Description for Genotyping

Objects from the Class

Objects of this class are unions of Quantification and Calls.

Slots

```
assayData: Object of class "AssayData"
phenoData: Object of class "AnnotatedDataFrame"
featureData: Object of class "AnnotatedDataFrame"
experimentData: Object of class "MIAME"
annotation: Object of class "character"
.__classVersion__: Object of class "Versions"
```

Extends

Class "SnpQSet", directly. Class "SnpCnvQSet", directly. Class "SnpCallSet", directly. Class "QuantificationSet", by class "SnpQSet", distance 2. Class eSet, by class "SnpQSet", distance 3. Class VersionedBiobase, by class "SnpQSet", distance 4. Class Versioned, by class "SnpQSet", distance 5. Class "QuantificationSet", by class "SnpCnvQSet", distance 2. Class eSet, by class "SnpCnvQSet", distance 3. Class VersionedBiobase, by class "SnpCnvQSet", distance 4. Class Versioned, by class "SnpCnvQSet", distance 5. Class "SnpLevelSet", by class "SnpCallSet", distance 2. Class eSet, by class "SnpCallSet", distance 3. Class VersionedBiobase, by class "SnpCallSet", distance 4. Class Versioned, by class "SnpCallSet", distance 5.

Methods

```
db signature(object="SnpCallSetPlus")
```

Examples

```
showClass("SnpCallSetPlus")
```

SnpCnvQSet-class *Class "SnpCnvQSet"*

Description

A class to store summarized measures from SnpCnvFeatureSet objects

Objects from the Class

Objects can be created by calls of the form `new("SnpCnvQSet", assayData, phenoData, featureData, experimentData, annotation, ...)`.

Slots

```
assayData: Object of class "AssayData"
phenoData: Object of class "AnnotatedDataFrame"
featureData: Object of class "AnnotatedDataFrame"
experimentData: Object of class "MIAME"
annotation: Object of class "character"
.__classVersion__: Object of class "Versions"
```

Extends

Class `QuantificationSet`, directly. Class `eSet`, by class "QuantificationSet", distance 2. Class `VersionedBiobase`, by class "QuantificationSet", distance 3. Class `Versioned`, by class "QuantificationSet", distance 4.

Methods

No methods defined with class "SnpCnvQSet" in the signature.

Examples

```
showClass("SnpCnvQSet")
```

cnConfidence	<i>Accessors for SnpCopyNumberSet objects</i>
--------------	---

Description

Accessors and replacement methods for SnpCopyNumberSet objects.

Usage

```
copyNumber(object)
cnConfidence(object)
copyNumber(object) <-value
cnConfidence(object) <-value
```

Arguments

object	SnpCopyNumberSet object
value	matrix

Value

Matrix with R rows (SNP's) and C columns (Samples) containing copy number estimates and confidences.

SnpLevelSet-class *Class "SnpLevelSet"*

Description

Virtual class for SNP-level data

Objects from the Class

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

Slots

assayData: The usual restrictions on assayData: contains matrices with equal dimensions, and with column number equal to nrow(phenoData). Class:[class.AssayData](#)

phenoData: See [class.eSet](#)

featureData: See [class.eSet](#)

experimentData: See [class.eSet](#)

annotation: See [class.eSet](#)

.__classVersion__: Object of class "Versions"

Extends

Class `eSet`, directly. Class `VersionedBiobase`, by class "eSet", distance 2. Class `Versioned`, by class "eSet", distance 3.

Methods

calls signature(object = "SnpLevelSet"): Accessor for matrix of genotype calls

calls<- signature(object = "SnpLevelSet", value = "matrix"): Replacement method for genotype calls.

callsConfidence signature(object = "SnpLevelSet"): Accessor for matrix of confidence scores for the genotype calls

callsConfidence<- signature(object = "SnpLevelSet", value = "matrix"): Replacement method

chromosome signature(object = "SnpLevelSet"): See also [chromosome](#)

chromosome<- signature(object = "SnpLevelSet", value = "character"): See also [chromosome](#)

cnConfidence signature(object = "SnpLevelSet"): Accessor for matrix of confidence scores for the copy number estimates.

cnConfidence<- signature(object = "SnpLevelSet", value = "matrix"): Replacement method for confidence scores of copy number estimates.

combine signature(x = "SnpLevelSet", y = "SnpLevelSet"): Useful for combining two objects with the same phenoData. For instance, combine an object created from the Hind 50k chip and an object created from the Xba 50k chip.

copyNumber signature(object = "SnpLevelSet"): Accessor for matrix of copy numbers.

copyNumber<- signature(object = "SnpLevelSet", value = "matrix"): Replacement method for copy number matrices

db signature(object = "SnpLevelSet"): See also [db](#)

position signature(object = "SnpLevelSet"): See also [position](#)

Author(s)

RS

Examples

```
showClass("SnpLevelSet")
```

SnpQSet-class

Class "SnpQSet"

Description

Class to store Snp (Quantifications) Summaries

Objects from the Class

Objects can be created by calls of the form `new("SnpQSet", assayData, senseThetaA, senseThetaB, antisenseThetaA, antisenseThetaB, phenoData, featureData, experimentData, annotation)`.

Slots

`assayData`: Object of class "AssayData"
`phenoData`: Object of class "AnnotatedDataFrame"
`featureData`: Object of class "AnnotatedDataFrame"
`experimentData`: Object of class "MIAME"
`annotation`: Object of class "character"
`.__classVersion__`: Object of class "Versions"

Extends

Class `eSet`, directly. Class `VersionedBiobase`, by class "eSet", distance 2. Class `Versioned`, by class "eSet", distance 3.

Methods

No methods defined with class "SnpQSet" in the signature.

See Also

[eSet](#), [VersionedBiobase](#), [Versioned](#)

getA

Compute average log-intensities / log-ratios

Description

Methods to compute average log-intensities and log-ratios across alleles, within strand.

Usage

```
getA(object)
getM(object)
```

Arguments

`object` SnpQSet, SnpCnvQSet or TilingFeatureSet2 object.

Details

For SNP data, SNPRMA summarizes the SNP information into 4 quantities (log2-scale):

- antisenseThetaAantisense allele A
- antisenseThetaBantisense allele B
- senseThetaAsense allele A
- senseThataBsense allele B

The average log-intensities are given by: $(\text{antisenseThetaA} + \text{antisenseThetaB}) / 2$ and $(\text{senseThetaA} + \text{senseThetaB}) / 2$.

The average log-ratios are given by: $\text{antisenseThetaA} - \text{antisenseThetaB}$ and $\text{senseThetaA} - \text{senseThetaB}$.

For Tiling data, `getM` and `getA` return the log-ratio and average log-intensities computed across channels: $M = \log_2(\text{channel1}) - \log_2(\text{channel2})$ $A = (\log_2(\text{channel1}) + \log_2(\text{channel2})) / 2$

Value

A 3-dimensional array (SNP's x Samples x Strand) with the requested measure, when the input SNP data (50K, 250K).

A 2-dimensional array (SNP's x Samples), when the input is from SNP 5.0 and SNP 6.0 arrays.

A 2-dimensional array if the input is from Tiling arrays.

See Also

[snprma](#)

SnpCallSet-class *"SnpSet" Extensions*

Description

Class to store genotype calls / copy number estimates (and confidences) obtained via SNP arrays.

Objects from the Class

Objects can be created by calls of the form `new("SnpCallSet", assayData, featureData, phenoData, experimentData, annotation, calls, callsConfidence, ...)`.

Slots

`assayData`: Object of class "AssayData"
`phenoData`: Object of class "AnnotatedDataFrame"
`featureData`: Object of class "AnnotatedDataFrame"
`experimentData`: Object of class "MIAME"
`annotation`: Object of class "character"
`.__classVersion__`: Object of class "Versions"

Extends

Class `eSet`, directly. Class `VersionedBiobase`, by class "eSet", distance 2. Class `Versioned`, by class "eSet", distance 3.

Methods

No methods defined with class "SnpCallSet" in the signature.

See Also

`eSet`, `VersionedBiobase`, `Versioned`

<code>affyPlatforms</code>	<i>Available Affymetrix platforms for SNP arrays</i>
----------------------------	--

Description

Provides a listing of available Affymetrix platforms currently supported by the R package `oligo`

Usage

```
affyPlatforms ()
```

Value

A vector of class character.

Author(s)

R. Scharpf

Examples

```
affyPlatforms ()
```

<code>chromosome</code>	<i>Accessor for chromosome information</i>
-------------------------	--

Description

Returns chromosome information.

Usage

```
pmChr (object)
```

Arguments

`object` `TilingFeatureSet` or `SnpCallSet` object

Details

`chromosome()` returns the chromosomal information for all probes and `pmChr()` subsets the output to the PM probes only (if a `TilingFeatureSet` object).

Value

Vector with chromosome information.

<code>efsExample</code>	<i>ExpressionFeatureSet Object</i>
-------------------------	------------------------------------

Description

Example of `ExpressionFeatureSet` Object.

Usage

```
data(efsExample)
```

Format

Object belongs to `ExpressionFeatureSet` class.

Examples

```
data(efsExample)
class(efsExample)
```

<code>scqsExample</code>	<i>SnpCnvQSet Example</i>
--------------------------	---------------------------

Description

Example of `SnpCnvQSet` object.

Usage

```
data(scqsExample)
```

Format

Object belongs to `SnpCnvQSet` class.

Examples

```
data(scqsExample)
class(scqsExample)
```

`sfsExample`*SnpFeatureSet Example*

Description

Example of SnpFeatureSet object.

Usage

```
data(sfsExample)
```

Format

Object belongs to SnpFeatureSet class

Examples

```
data(sfsExample)
class(sfsExample)
```

`sqsExample`*SnpQSet Example*

Description

Example of SnpQSet instance.

Usage

```
data(sqsExample)
```

Format

Belongs to SnpQSet class.

Examples

```
data(sqsExample)
class(sqsExample)
```

 db

Get the connection to the SQLite Database

Description

This function will return the SQLite connection to the database associated to objects used in oligo.

Usage

```
db(object)
```

Arguments

`object` Object of valid class. See methods.

Value

SQLite connection.

Methods

object = "FeatureSet" object of class FeatureSet
object = "SnpCallSet" object of class SnpCallSet
object = "DBPDInfo" object of class DBPDInfo
object = "SnpLevelSet" object of class SnpLevelSet

Author(s)

Benilton Carvalho

Examples

```
## db(object)
```

 exprs-methods

Accessor for the 'exprs' slot

Description

Accessor for the 'exprs'/se.exprs' slot of FeatureSet-like objects

Methods

object = "ExpressionSet" Expression matrix for objects of this class. Usually results of preprocessing algorithms, like RMA.
object = "FeatureSet" General container 'exprs' inherited from eSet
object = "SnpSet" General container 'exprs' inherited from eSet, not yet used.

genomeBuild	<i>Genome Build Information</i>
-------------	---------------------------------

Description

Returns the genome build information. This information comes from the annotation package and is given as an argument during the package creation process.

Usage

```
genomeBuild(object)
```

Arguments

object PDInfo or FeatureSet object.

getPlatformDesign	<i>Retrieve Platform Design object</i>
-------------------	--

Description

Retrieve platform design object.

Usage

```
getPlatformDesign(object)  
getPD(object)
```

Arguments

object FeatureSet object

Details

Retrieve platform design object.

Value

platformDesign or PDInfo object.

kind *Array type*

Description

Retrieves the array type.

Usage

```
kind(object)
```

Arguments

object FeatureSet or DBPDInfo object

Value

String: "Expression", "Exon", "SNP" or "Tiling"

Examples

```
if (require(pd.mapping50k.xba240)) {
  data(sfsExample)
  annotation(sfsExample) <- "pd.mapping50k.xba240"
  kind(sfsExample)
}
```

length-methods *Number of samples for FeatureSet-like objects.*

Description

Number of samples for FeatureSet-like objects.

Methods

x = "FeatureSet" Number of samples

manufacturer-methods *Manufacturer ID for FeatureSet-like objects.*

Description

Manufacturer ID for FeatureSet-like and DBPDInfo-like objects.

Methods

object = "FeatureSet" Manufacturer ID

object = "PDInfo" Manufacturer ID

`oligoSnpSet-methods`*Methods for oligoSnpSet class*

Description

Methods for oligoSnpSet

`pdPkgFromBioC`*Get packages from BioConductor.*

Description

This function checks if a given package is available on BioConductor and installs it, in case it is.

Usage

```
pdPkgFromBioC(pkgname, lib = .libPaths()[1], verbose = TRUE)
```

Arguments

<code>pkgname</code>	character. Name of the package to be installed.
<code>lib</code>	character. Path where to install the package at.
<code>verbose</code>	logical. Verbosity flag.

Details

Internet connection required.

Value

Logical: TRUE if package was found, downloaded and installed; FALSE otherwise.

Author(s)

Benilton Carvalho

See Also

`download.packages`

Examples

```
## Not run:
pdPkgFromBioC("pd.mapping50k.xba240")

## End(Not run)
```

platform-methods *Platform Information*

Description

Platform Information

Methods

object = "FeatureSet" platform information

pmAllele *Access the allele information for PM probes.*

Description

Accessor to the allelic information for PM probes.

Usage

pmAllele(object)

Arguments

object SnpFeatureSet or PDInfo object.

pmFragmentLength-methods
Information on Fragment Length

Description

This method will return the fragment length for PM probes.

Methods

object = "AffySNPPDInfo" On AffySNPPDInfo objects, it will return the fragment length that contains the SNP in question.

pmPosition	<i>Accessor to position information</i>
------------	---

Description

pmPosition will return the genomic position for the (PM) probes.

Usage

```
pmPosition(object)  
pmOffset(object)
```

Arguments

object AffySNPPDInfo, TilingFeatureSet or SnpCallSet object

Details

pmPosition will return genomic position for PM probes on a tiling array.

pmOffset will return the offset information for PM probes on SNP arrays.

pmStrand	<i>Accessor to the strand information</i>
----------	---

Description

Returns the strand information on SNP arrays for PM probes (0 - sense / 1 - antisense).

Usage

```
pmStrand(object)
```

Arguments

object AffySNPPDInfo object

position	<i>Accessor to position information</i>
----------	---

Description

position will return the genomic position of a SNP.

Usage

```
position(object)
```

Arguments

object object inheriting from SnpLevelSet

Details

position will return genomic position of a SNP (number of basepairs from the 5-prime chromosomal end)

Value

an integer

Author(s)

R. Scharpf

probeNames	<i>Accessor to feature names</i>
------------	----------------------------------

Description

Accessor to PM feature names.

Usage

```
probeNames(object, subset = NULL)
```

Arguments

object FeatureSet or DBPDInfo
subset not implemented yet.

requireAnnotation *Helper function to load packages.*

Description

This function checks the existence of a given package and loads it if available. If the package is not available, the function checks its availability on BioConductor, downloads it and installs it.

Usage

```
requireAnnotation(pkgname, lib=.libPaths()[1], verbose = TRUE)
```

Arguments

pkgname	character. Package name (usually an annotation package).
lib	character. Path where to install packages at.
verbose	logical. Verbosity flag.

Value

Logical: TRUE if package is available or FALSE if package unavailable for download.

Author(s)

Benilton Carvalho

See Also

install.packages

Examples

```
## Not run:
requirePackage("pd.mapping50k.xba240")

## End (Not run)
```

sampleNames-methods

Sample names for FeatureSet-like objects

Description

Returns sample names for FeatureSet-like objects.

Methods

object = "FeatureSet" Sample names

antisenseThetaA *Summarized Measures for SNP*

Description

SNPRMA summarizes the intensities for each SNP into 4 quantities. These measures are at the log₂ scale and are used to create log-ratios and log-averages, which are used by the genotype algorithm and copy number tool.

Usage

```
antisenseThetaA(object)
antisenseThetaB(object)
senseThetaA(object)
senseThetaB(object)
thetaA(object)
thetaB(object)
```

Arguments

object SnpQSet object

Details

For a given SNP, an Affymetrix SNP array usually contains multiple features for each combination allele x strand, on which the summaries are computed.

Value

A matrix with R rows (SNP's) x C columns (Samples).

See Also

[snprma](#), [crlmm](#)

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