

# Package: BiocGenerics (via r-universe)

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**Title** S4 generic functions used in Bioconductor

**Description** The package defines many S4 generic functions used in Bioconductor.

**biocViews** Infrastructure

**URL** <https://bioconductor.org/packages/BiocGenerics>

**BugReports** <https://github.com/Bioconductor/BiocGenerics/issues>

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**Collate** S3-classes-as-S4-classes.R utils.R normarg-utils.R  
replaceSlots.R aperm.R append.R as.data.frame.R as.list.R  
as.vector.R cbind.R do.call.R duplicated.R eval.R Extremes.R  
format.R funprog.R get.R grep.R is.unsorted.R lapply.R mapply.R  
match.R mean.R nrow.R order.R paste.R rank.R rep.R  
row\_colnames.R saveRDS.R sets.R sort.R start.R subset.R t.R  
table.R tapply.R unique.R unlist.R unsplit.R relist.R var.R  
which.R which.min.R boxplot.R image.R density.R IQR.R mad.R  
residuals.R weights.R xtabs.R annotation.R combine.R  
containsOutOfMemoryData.R dbconn.R dge.R dims.R fileName.R  
normalize.R Ontology.R organism\_species.R path.R plotMA.R  
plotPCA.R score.R strand.R toTable.R type.R updateObject.R  
testPackage.R zzz.R

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BiocGenerics-package    *S4 generic functions for Bioconductor*

---

## Description

S4 generic functions needed by many Bioconductor packages.

## Details

We divide the generic functions defined in the **BiocGenerics** package in 2 categories: (1) functions already defined in base R and explicitly promoted to generics in BiocGenerics, and (2) Bioconductor specific generics.

### (1) Functions defined in base R and explicitly promoted to generics in the BiocGenerics package:

Generics for functions defined in package **base**:

- BiocGenerics::aperm
- BiocGenerics::append
- BiocGenerics::as.data.frame
- BiocGenerics::as.list
- BiocGenerics::as.vector
- BiocGenerics::rbind, BiocGenerics::cbind
- BiocGenerics::do.call
- BiocGenerics::duplicated, BiocGenerics::anyDuplicated
- BiocGenerics::eval
- Extremes: BiocGenerics::pmax, BiocGenerics::pmin, BiocGenerics::pmax.int, BiocGenerics::pmin.int
- BiocGenerics::format
- funprog: BiocGenerics::Reduce, BiocGenerics::Filter, BiocGenerics::Find, BiocGenerics::Map, BiocGenerics::Position
- BiocGenerics::get, BiocGenerics::mget
- BiocGenerics::grep, BiocGenerics::grepl
- BiocGenerics::is.unsorted
- BiocGenerics::lapply, BiocGenerics::sapply
- BiocGenerics::mapply
- BiocGenerics::match, BiocGenerics::%in%
- BiocGenerics::nrow, BiocGenerics::ncol, BiocGenerics::NROW, BiocGenerics::NCOL
- BiocGenerics::order
- BiocGenerics::paste
- BiocGenerics::rank
- BiocGenerics::rep.int
- BiocGenerics::rownames, BiocGenerics::rownames<-, BiocGenerics::colnames, BiocGenerics::colnames<-
- BiocGenerics::saveRDS
- sets: BiocGenerics::union, BiocGenerics::intersect, BiocGenerics::setdiff
- BiocGenerics::sort
- BiocGenerics::start, BiocGenerics::start<-, BiocGenerics::end, BiocGenerics::end<-, BiocGenerics::width, BiocGenerics::width<-, BiocGenerics::pos
- BiocGenerics::subset
- BiocGenerics::t
- BiocGenerics::table
- BiocGenerics::tapply
- BiocGenerics::unique
- BiocGenerics::unlist
- BiocGenerics::unsplit
- BiocGenerics::which
- BiocGenerics::which.min, BiocGenerics::which.max

Generics for functions defined in package **utils**:

- BiocGenerics::relist

Generics for functions defined in package **graphics**:

- `BiocGenerics::boxplot`
- `BiocGenerics::image`

Generics for functions defined in package **stats**:

- `BiocGenerics::density`
- `BiocGenerics::residuals`
- `BiocGenerics::weights`
- `BiocGenerics::xtabs`

**(2) Bioconductor specific generics:**

- `annotation, annotation<-`
- `combine`
- `containsOutOfMemoryData`
- `dbconn, dbfile`
- `counts, counts<-, design, design<-, dispTable, dispTable<-, sizeFactors, sizeFactors<-, conditions, conditions<-, estimateSizeFactors, estimateDispersions, plotDispEsts`
- `dims, nrows, ncols,`
- `fileName`
- `normalize`
- `Ontology`
- `organism, organism<-, species, species<-`
- `path, path<-, basename, basename<-, dirname, dirname<-`
- `plotMA`
- `plotPCA`
- `score, score<-`
- `strand, strand<-, invertStrand`
- `toTable`
- `type, type<-`
- `updateObject`

**Note**

More generics can be added on request by sending an email to the Bioc-devel mailing list:

<http://bioconductor.org/help/mailling-list/>

Things that should NOT be added to the **BiocGenerics** package:

- Internal generic primitive functions like `length`, `dim`, ``dim<-``, etc... See `?InternalMethods` for the complete list. There are a few exceptions though, that is, the **BiocGenerics** package may actually redefine a few of those internal generic primitive functions as S4 generics when for example the signature of the internal generic primitive is not appropriate (this is the case for `BiocGenerics::cbind`).
- S3 and S4 group generic functions like `Math`, `Ops`, etc... See `?groupGeneric` and `?S4groupGeneric` for the complete list.
- Generics already defined in the **stats4** package.

**Author(s)**

The Bioconductor Dev Team

**See Also**

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [setGeneric](#) and [setMethod](#) for defining generics and methods.

**Examples**

```
## List all the symbols defined in this package:
ls('package:BiocGenerics')
```

---

annotation

*Accessing annotation information*

---

**Description**

Get or set the annotation information contained in an object.

**Usage**

```
annotation(object, ...)
annotation(object, ...) <- value
```

**Arguments**

object	An object containing annotation information.
...	Additional arguments, for use in specific methods.
value	The annotation information to set on object.

**See Also**

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [annotation,eSet-method](#) in the **Biobase** package for an example of a specific annotation method (defined for [eSet](#) objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
annotation
showMethods("annotation")

library(Biobase)
showMethods("annotation")
selectMethod("annotation", "eSet")
```

---

aperm

*Transposing an array-like object*

---

## Description

Transpose an array-like object by permuting its dimensions.

This is a multidimensional generalization of the `t()` operator used for 2D-transposition.

NOTE: This man page is for the `aperm` *S4 generic function* defined in the **BiocGenerics** package. See `?base::aperm` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
aperm(a, perm, ...)
```

## Arguments

<code>a</code>	An array-like object.
<code>perm, ...</code>	See <code>?base::aperm</code> for a description of these arguments.

## Value

A transposed version of array-like object `a`, with subscripts permuted as indicated by the `perm` vector.

## See Also

- `base::aperm` for the default `aperm` method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `aperm,SVT_SparseArray-method` in the **SparseArray** package for an example of a specific `aperm` method (defined for `SVT_SparseArray` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
aperm # note the dispatch on the 'a' arg only
showMethods("aperm")
selectMethod("aperm", "ANY") # the default method
```

---

append

*Append elements to a vector-like object*

---

### Description

Append (or insert) elements to (in) a vector-like object.

NOTE: This man page is for the `append` *S4 generic function* defined in the **BiocGenerics** package. See `?base::append` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-like or data-frame-like) not supported by the default method.

### Usage

```
append(x, values, after=length(x))
```

### Arguments

<code>x</code>	The vector-like object to be modified.
<code>values</code>	The vector-like object containing the values to be appended to <code>x</code> . <code>values</code> would typically be of the same class as <code>x</code> , but not necessarily.
<code>after</code>	A subscript, after which the values are to be appended.

### Value

See `?base::append` for the value returned by the default method.

Specific methods defined in Bioconductor packages will typically return an object of the same class as `x` and of length `length(x) + length(values)`.

### See Also

- `base::append` for the default append method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `append,Vector,Vector-method` in the **S4Vectors** package for an example of a specific append method (defined for `Vector` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

### Examples

```
append # note the dispatch on the 'x' and 'values' args only
showMethods("append")
selectMethod("append", c("ANY", "ANY")) # the default method
```

---

as.data.frame	<i>Coerce to a data frame</i>
---------------	-------------------------------

---

## Description

Generic function to coerce to a data frame, if possible.

NOTE: This man page is for the `as.data.frame` *S4 generic function* defined in the **BiocGenerics** package. See `?base::as.data.frame` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
as.data.frame(x, row.names=NULL, optional=FALSE, ...)
```

## Arguments

`x` The object to coerce.

`row.names`, `optional`, ...

See `?base::as.data.frame` for a description of these arguments.

## Value

An ordinary data frame.

See `?base::as.data.frame` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## See Also

- `base::as.data.frame` for the default `as.data.frame` method.
- `toTable` for an alternative to `as.data.frame`.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `as.data.frame,DataFrame-method` in the **S4Vectors** package, and `as.data.frame,IntegerRanges-method` in the **IRanges** package, for examples of specific `as.data.frame` methods (defined for `DataFrame` and `IntegerRanges` objects, respectively).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
as.data.frame # note the dispatch on the 'x' arg only
showMethods("as.data.frame")
selectMethod("as.data.frame", "ANY") # the default method
```

---

`as.list`*Coerce to a list*

---

### Description

Generic function to coerce to a list, if possible.

NOTE: This man page is for the `as.list` *S4 generic function* defined in the **BiocGenerics** package. See `?base::as.list` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects not supported by the default method.

### Usage

```
as.list(x, ...)
```

### Arguments

<code>x</code>	The object to coerce.
<code>...</code>	Additional arguments, for use in specific methods.

### Value

An ordinary list.

### See Also

- `base::as.list` for the default `as.list` method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `as.list,List-method` in the **S4Vectors** package for an example of a specific `as.list` method (defined for `List` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

### Examples

```
as.list
showMethods("as.list")
selectMethod("as.list", "ANY") # the default method

library(S4Vectors)
showMethods("as.list")
## The as.list() method for List objects:
selectMethod("as.list", "List")
```

---

as.vector	<i>Coerce an object into a vector</i>
-----------	---------------------------------------

---

## Description

Attempt to coerce an object into a vector of the specified mode. If the mode is not specified, attempt to coerce to whichever vector mode is considered more appropriate for the class of the supplied object.

NOTE: This man page is for the `as.vector` *S4 generic function* defined in the **BiocGenerics** package. See `?base::as.vector` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
as.vector(x, mode="any")
```

## Arguments

x	The object to coerce.
mode	See <code>?base::as.vector</code> for a description of this argument.

## Value

A vector.

See `?base::as.vector` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## See Also

- `base::as.vector` for the default `as.vector` method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `as.vector,Rle-method` in the **S4Vectors** package, and `as.vector,AtomicList-method` in the **IRanges** packages, for examples of specific `as.vector` methods (defined for **Rle** and **AtomicList** objects, respectively).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
as.vector # note the dispatch on the 'x' arg only
showMethods("as.vector")
selectMethod("as.vector", "ANY") # the default method
```

---

boxplot

*Box plots*

---

## Description

Produce box-and-whisker plot(s) of the given (grouped) values.

NOTE: This man page is for the `boxplot` *S4 generic function* defined in the **BiocGenerics** package. See `?graphics::boxplot` for the default method (defined in the **graphics** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
boxplot(x, ...)
```

## Arguments

`x, ...` See `?graphics::boxplot`.

## Value

See `?graphics::boxplot` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## See Also

- `graphics::boxplot` for the default boxplot method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `boxplot,AffyBatch-method` in the **affy** package for an example of a specific boxplot method (defined for `AffyBatch` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
boxplot
showMethods("boxplot")
selectMethod("boxplot", "ANY") # the default method

library(affy)
showMethods("boxplot")
## The boxplot() method for AffyBatch objects:
selectMethod("boxplot", "AffyBatch")
```

---

`cbind`*Combine objects by rows or columns*

---

## Description

`rbind` and `cbind` take one or more objects and combine them by columns or rows, respectively.

NOTE: This man page is for the `rbind` and `cbind` *S4 generic functions* defined in the **BiocGenerics** package. See `?base::cbind` for the default methods (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-like or matrix-like) not supported by the default methods.

## Usage

```
rbind(..., deparse.level=1)
cbind(..., deparse.level=1)
```

## Arguments

`...` One or more vector-like or matrix-like objects. These can be given as named arguments.

`deparse.level` See `?base::cbind` for a description of this argument.

## Value

See `?base::cbind` for the value returned by the default methods.

Specific methods defined in Bioconductor packages will typically return an object of the same class as the input objects.

## See Also

- `base::cbind` for the default `rbind` and `cbind` methods.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `rbind,RectangularData-method` and `cbind,DataFrame-method` in the **S4Vectors** package for examples of specific `rbind` and `cbind` methods (defined for `RectangularData` derivatives and `DataFrame` objects, respectively).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
rbind # note the dispatch on the '...' arg only
showMethods("rbind")
selectMethod("rbind", "ANY") # the default method

cbind # note the dispatch on the '...' arg only
showMethods("cbind")
```

```

selectMethod("cbind", "ANY") # the default method

library(S4Vectors)
showMethods("rbind")
## The rbind() method for RectangularData derivatives:
selectMethod("rbind", "RectangularData")
## The cbind() method for DataFrame objects:
selectMethod("cbind", "DataFrame")

```

---

combine

*Combining or merging different Bioconductor data structures*


---

### Description

The combine generic function handles methods for combining or merging different Bioconductor data structures. It should, given an arbitrary number of arguments of the same class (possibly by inheritance), combine them into a single instance in a sensible way (some methods may only combine 2 objects, ignoring ... in the argument list; because Bioconductor data structures are complicated, check carefully that combine does as you intend).

### Usage

```
combine(x, y, ...)
```

### Arguments

x	One of the values.
y	A second value.
...	Any other objects of the same class as x and y.

### Details

There are two basic combine strategies. One is an intersection strategy. The returned value should only have rows (or columns) that are found in all input data objects. The union strategy says that the return value will have all rows (or columns) found in any one of the input data objects (in which case some indication of what to use for missing values will need to be provided).

These functions and methods are currently under construction. Please let us know if there are features that you require.

### Value

A single value of the same class as the most specific common ancestor (in class terms) of the input values. This will contain the appropriate combination of the data in the input values.

## Methods

The following methods are defined in the **BiocGenerics** package:

`combine(x=ANY, missing)` Return the first (x) argument unchanged.

`combine(data.frame, data.frame)` Combines two `data.frame` objects so that the resulting `data.frame` contains all rows and columns of the original objects. Rows and columns in the returned value are unique, that is, a row or column represented in both arguments is represented only once in the result. To perform this operation, `combine` makes sure that data in shared rows and columns are identical in the two `data.frames`. Data differences in shared rows and columns usually cause an error. `combine` issues a warning when a column is a `factor` and the levels of the factor in the two `data.frames` are different.

`combine(matrix, matrix)` Combined two `matrix` objects so that the resulting `matrix` contains all rows and columns of the original objects. Both matrices must have `dimnames`. Rows and columns in the returned value are unique, that is, a row or column represented in both arguments is represented only once in the result. To perform this operation, `combine` makes sure that data in shared rows and columns are all equal in the two matrices.

Additional `combine` methods are defined in the **Biobase** package for `AnnotatedDataFrame`, `AssayData`, `MIAME`, and `eSet` objects.

## Author(s)

Biocore

## See Also

- [combine,AnnotatedDataFrame,AnnotatedDataFrame-method](#), [combine,AssayData,AssayData-method](#), [combine,MIAME,MIAME-method](#), and [combine,eSet,eSet-method](#) in the **Biobase** package for additional `combine` methods.
- [merge](#) for merging two data frames (or data-frame-like) objects.
- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
combine
showMethods("combine")
selectMethod("combine", c("ANY", "missing"))
selectMethod("combine", c("data.frame", "data.frame"))
selectMethod("combine", c("matrix", "matrix"))

## -----
## COMBINING TWO DATA FRAMES
## -----
x <- data.frame(x=1:5,
                y=factor(letters[1:5], levels=letters[1:8]),
                row.names=letters[1:5])
```

```

y <- data.frame(z=3:7,
               y=factor(letters[3:7], levels=letters[1:8]),
               row.names=letters[3:7])
combine(x,y)

w <- data.frame(w=4:8,
               y=factor(letters[4:8], levels=letters[1:8]),
               row.names=letters[4:8])
combine(w, x, y)

# y is converted to 'factor' with different levels
df1 <- data.frame(x=1:5,y=letters[1:5], row.names=letters[1:5])
df2 <- data.frame(z=3:7,y=letters[3:7], row.names=letters[3:7])
try(combine(df1, df2)) # fails
# solution 1: ensure identical levels
y1 <- factor(letters[1:5], levels=letters[1:7])
y2 <- factor(letters[3:7], levels=letters[1:7])
df1 <- data.frame(x=1:5,y=y1, row.names=letters[1:5])
df2 <- data.frame(z=3:7,y=y2, row.names=letters[3:7])
combine(df1, df2)
# solution 2: force column to be 'character'
df1 <- data.frame(x=1:5,y=I(letters[1:5]), row.names=letters[1:5])
df2 <- data.frame(z=3:7,y=I(letters[3:7]), row.names=letters[3:7])
combine(df1, df2)

## -----
## COMBINING TWO MATRICES
## -----
m <- matrix(1:20, nrow=5, dimnames=list(LETTERS[1:5], letters[1:4]))
combine(m[1:3,], m[4:5,])
combine(m[1:3, 1:3], m[3:5, 3:4]) # overlap

```

---

containsOutOfMemoryData

*Does an object contain out-of-memory data?*

---

## Description

Some objects in Bioconductor can use on-disk or other out-of-memory representation for their data, typically (but not necessarily) when the data is too big to fit in memory. For example the data in a [TxDb](#) object is stored in an SQLite database, and the data in an [HDF5Array](#) object is stored in an HDF5 file.

The `containsOutOfMemoryData()` function determines whether an object contains out-of-memory data or not.

Note that objects with out-of-memory data are usually not compatible with a serialization/unserialization roundtrip. More concretely, base: `:saveRDS()/base: :readRDS()` tend to silently break them!

See [?saveHDF5SummarizedExperiment](#) in the **HDF5Array** package for a more extensive discussion about this.

## Usage

```
containsOutOfMemoryData(object)
```

## Arguments

object            The object to be tested.

## Details

An object can store *some* of its data on disk and *some* of it in memory. This is the case for example when a [SummarizedExperiment](#) object (or derivative) has some of its assays on disk (e.g. in [HDF5Matrix](#) objects) and others in memory (e.g. in ordinary matrices and/or [SparseMatrix](#) objects).

Of course in this case, `containsOutOfMemoryData()` will still return TRUE. In other words, `containsOutOfMemoryData(object)` will only return FALSE when all the data in object resides in memory, that is, when the object can safely be serialized.

## Value

TRUE or FALSE.

## Note

TO DEVELOPERS:

The **BiocGenerics** package also defines the following:

- A default `containsOutOfMemoryData()` method that returns TRUE if object is an S4 object with at least one slot for which `containsOutOfMemoryData()` is TRUE (recursive definition), and FALSE otherwise.
- A `containsOutOfMemoryData()` method for list objects that returns TRUE if object has at least one list element for which `containsOutOfMemoryData()` is TRUE (recursive definition), and FALSE otherwise.
- A `containsOutOfMemoryData()` method for environment objects that returns TRUE if object contains at least one object for which `containsOutOfMemoryData()` is TRUE (recursive definition), and FALSE otherwise.
- The `OutOfMemoryObject` class. This is a virtual S4 class with no slots that any class defined in Bioconductor that represents out-of-memory objects should extend.
- A `containsOutOfMemoryData()` method for `OutOfMemoryObject` derivatives that returns TRUE.

Therefore, if you implement a class that uses an out-of-memory representation, make sure that it contains the `OutOfMemoryObject` class. This will make `containsOutOfMemoryData()` return TRUE on your objects, so you don't need to define a `containsOutOfMemoryData()` method for them.

## See Also

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```

containsOutOfMemoryData
showMethods("containsOutOfMemoryData")

## The default method:
selectMethod("containsOutOfMemoryData", "ANY")

## The method for list objects:
selectMethod("containsOutOfMemoryData", "list")

## The method for OutOfMemoryObject derivatives:
selectMethod("containsOutOfMemoryData", "OutOfMemoryObject")

m <- matrix(0, nrow=7, ncol=10)
m[sample(length(m), 20)] <- runif(20)
containsOutOfMemoryData(m) # FALSE

library(SparseArray)
svt <- as(m, "SparseArray")
svt
containsOutOfMemoryData(m) # FALSE
containsOutOfMemoryData(list(m, svt)) # FALSE

library(HDF5Array)
M <- as(m, "HDF5Array")
M
containsOutOfMemoryData(M) # TRUE
containsOutOfMemoryData(list(m, svt, M)) # TRUE

```

---

dbconn

*Accessing SQLite DB information*


---

**Description**

Get a connection object or file path for a SQLite DB

**Usage**

```

dbconn(x)
dbfile(x)

```

**Arguments**

x                    An object with a SQLite connection.

**Value**

dbconn returns a connection object to the SQLite DB containing x's data.

dbfile returns a path (character string) to the SQLite DB (file) containing x's data.

**See Also**

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [dbconn,AnnotationDb-method](#) in the **AnnotationDbi** package for an example of a specific dbconn method (defined for [dbconn](#) objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
dbconn
showMethods("dbconn")
dbfile
showMethods("dbfile")

library(AnnotationDbi)
showMethods("dbconn")
selectMethod("dbconn", "AnnotationDb")
```

---

density

*Kernel density estimation*

---

**Description**

The generic function `density` computes kernel density estimates.

NOTE: This man page is for the density *S4 generic function* defined in the **BiocGenerics** package. See `?stats::density` for the default method (defined in the **stats** package). Bioconductor packages can define specific methods for objects not supported by the default method.

**Usage**

```
density(x, ...)
```

**Arguments**

`x, ...` See `?stats::density`.

**Value**

See `?stats::density` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

**See Also**

- `stats::density` for the default density method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `density,flowClust-method` in the **flowClust** package for an example of a specific density method (defined for **flowClust** objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
density
showMethods("density")
selectMethod("density", "ANY") # the default method
```

---

 dge

---

*Accessors and generic functions used in the context of count datasets*


---

**Description**

These generic functions provide basic interfaces to operations on and data access to count datasets.

**Usage**

```
counts(object, ...)
counts(object, ...) <- value
design(object, ...)
design(object, ...) <- value
dispTable(object, ...)
dispTable(object, ...) <- value
sizeFactors(object, ...)
sizeFactors(object, ...) <- value
conditions(object, ...)
conditions(object, ...) <- value
estimateSizeFactors(object, ...)
estimateDispersions(object, ...)
plotDispEsts(object, ...)
```

**Arguments**

<code>object</code>	Object of class for which methods are defined, e.g., <code>CountDataSet</code> , <code>DESeqSummarizedExperiment</code> or <code>ExonCountSet</code> .
<code>value</code>	Value to be assigned to corresponding components of <code>object</code> ; supported types depend on method implementation.
<code>...</code>	Further arguments, perhaps used by methods

## Details

For the details, please consult the manual pages of the methods in the **DESeq**, **DESeq2**, and **DEXSeq** packages and the package vignettes.

## Author(s)

W. Huber, S. Anders

## See Also

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

---

dims

*Get the dimensions of each element of a list-like object*

---

## Description

Get the dimensions, number of rows, or number of columns, of each element of a list-like object.

Note that these functions are the *vectorized versions* of corresponding functions `dim()`, `nrow()`, and `ncol()`, in the same fashion that `lengths()` is the *vectorized version* of `length`.

## Usage

```
dims(x, use.names=TRUE)
nrows(x, use.names=TRUE)
ncols(x, use.names=TRUE)
```

## Arguments

x	List-like object (or environment) where all the list elements are expected to be array-like objects with the <i>same number of dimensions</i> .
use.names	Logical indicating if the names on x should be propagated to the returned matrix (as its rownames) or vector (as its names).

## Value

For `dims()`: Typically a numeric matrix with one row per list element in x and one column per dimension in these list elements (they're all expected to have the same number of dimensions). The i-th row in the returned matrix is a vector containing the dimensions of the i-th list element in x. More formally:

```
dims(x)[i, ] is dim(x[[i]])
```

for any valid `i`. By default the names on `x`, if any, are propagated as the rownames of the returned matrix, unless `use.names` is set to `FALSE`.

For `nrows()` or `ncols()`: A numeric vector with one element per list element in `x`. The `i`-th element in the returned vector is the number of rows (or columns) of the `i`-th list element in `x`. More formally:

```
nrows(x)[i] is nrow(x[[i]]) and ncols(x)[i] is ncol(x[[i]])
```

for any valid `i`. By default the names on `x`, if any, are propagated on the returned vector, unless `use.names` is set to `FALSE`.

### See Also

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [dims,DataFrameList-method](#) in the **IRanges** package for an example of a specific `dims` method (defined for [DataFrameList](#) objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

### Examples

```
dims
showMethods("dims")

library(IRanges)
showMethods("dims")
selectMethod("dims", "DataFrameList")
```

---

do.call

*Execute a function call*

---

### Description

`do.call` constructs and executes a function call from a name or a function and a list of arguments to be passed to it.

NOTE: This man page is for the `do.call` *S4 generic function* defined in the **BiocGenerics** package. See `?base::do.call` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects not supported by the default method.

### Usage

```
do.call(what, args, quote=FALSE, envir=parent.frame())
```

**Arguments**

what	The default method expects either a function or a non-empty character string naming the function to be called. See <code>?base::do.call</code> for the details. Specific methods can support other objects. Please refer to the documentation of a particular method for the details.
args	The default method expects a <i>list</i> of arguments to the function call (the names attribute of <code>args</code> gives the argument names). See <code>?base::do.call</code> for the details. Specific methods can support other objects. Please refer to the documentation of a particular method for the details.
quote, envir	See <code>?base::do.call</code> for a description of these arguments.

**Value**

The result of the (evaluated) function call.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

**See Also**

- `base::do.call` for the default `do.call` method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `BiocGenerics` for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
do.call # note the dispatch on the 'what' and 'args' args only
showMethods("do.call")
selectMethod("do.call", c("ANY", "ANY")) # the default method
```

---

duplicated

*Determine duplicate elements*


---

**Description**

Determines which elements of a vector-like or data-frame-like object are duplicates of elements with smaller subscripts, and returns a logical vector indicating which elements (rows) are duplicates.

NOTE: This man page is for the `duplicated` and `anyDuplicated` *S4 generic functions* defined in the **BiocGenerics** package. See `?base::duplicated` for the default methods (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-like or data-frame-like) not supported by the default method.

**Usage**

```
duplicated(x, incomparables=FALSE, ...)
anyDuplicated(x, incomparables=FALSE, ...)
```

**Arguments**

`x` A vector-like or data-frame-like object.

`incomparables, ...` See `?base::duplicated` for a description of these arguments.

**Value**

The default `duplicated` method (see `?base::duplicated`) returns a logical vector of length `N` where `N` is:

- `length(x)` when `x` is a vector;
- `nrow(x)` when `x` is a data frame.

Specific `duplicated` methods defined in Bioconductor packages must also return a logical vector of the same length as `x` when `x` is a vector-like object, and a logical vector with one element for each row when `x` is a data-frame-like object.

The default `anyDuplicated` method (see `?base::duplicated`) returns a single non-negative integer and so must the specific `anyDuplicated` methods defined in Bioconductor packages.

`anyDuplicated` should always behave consistently with `duplicated`.

**See Also**

- `base::duplicated` for the default `duplicated` and `anyDuplicated` methods.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `duplicated,Rle-method` in the **S4Vectors** package for an example of a specific `duplicated` method (defined for **Rle** objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
duplicated
showMethods("duplicated")
selectMethod("duplicated", "ANY") # the default method

anyDuplicated
showMethods("anyDuplicated")
selectMethod("anyDuplicated", "ANY") # the default method
```

---

eval	<i>Evaluate an (unevaluated) expression</i>
------	---

---

## Description

eval evaluates an R expression in a specified environment.

NOTE: This man page is for the eval *S4 generic function* defined in the **BiocGenerics** package. See `?base::eval` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
eval(expr, envir=parent.frame(),
      enclos=if (is.list(envir) || is.pairlist(envir))
                parent.frame() else baseenv())
```

## Arguments

expr	An object to be evaluated. May be any object supported by the default method (see <code>?base::eval</code> ) or by the additional methods defined in Bioconductor packages.
envir	The <i>environment</i> in which expr is to be evaluated. May be any object supported by the default method (see <code>?base::eval</code> ) or by the additional methods defined in Bioconductor packages.
enclos	See <code>?base::eval</code> for a description of this argument.

## Value

See `?base::eval` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## See Also

- `base::eval` for the default eval method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `eval,expression,Vector-method` in the **IRanges** package for an example of a specific eval method (defined for when the expr and envir arguments are an `expression` and a `Vector` object, respectively).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
eval # note the dispatch on 'expr' and 'envir' args only
showMethods("eval")
selectMethod("eval", c("ANY", "ANY")) # the default method
```

---

evalq	<i>Evaluate an (unevaluated) expression</i>
-------	---

---

### Description

evalq evaluates an R expression (the quoted form of its first argument) in a specified environment.

NOTE: This man page is for the evalq wrapper defined in the **BiocGenerics** package. See `?base::evalq` for the function defined in the **base** package. This wrapper correctly delegates to the eval generic, rather than `base::eval`.

### Usage

```
evalq(expr, envir=parent.frame(),
      enclos=if (is.list(envir) || is.pairlist(envir))
                parent.frame() else baseenv())
```

### Arguments

expr	Quoted to form the expression that is evaluated.
envir	The <i>environment</i> in which expr is to be evaluated. May be any object supported by methods on the <code>eval</code> generic.
enclos	See <code>?base::evalq</code> for a description of this argument.

### Value

See `?base::evalq`.

### See Also

- `base::evalq` for the base evalq function.

### Examples

```
evalq # note just a copy of the original evalq
```

---

Extremes	<i>Maxima and minima</i>
----------	--------------------------

---

### Description

pmax, pmin, pmax.int and pmin.int return the parallel maxima and minima of the input values.

NOTE: This man page is for the pmax, pmin, pmax.int and pmin.int *S4 generic functions* defined in the **BiocGenerics** package. See `?base::pmax` for the default methods (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-like or matrix-like) not supported by the default methods.

## Usage

```
pmax(..., na.rm=FALSE)
pmin(..., na.rm=FALSE)

pmax.int(..., na.rm=FALSE)
pmin.int(..., na.rm=FALSE)
```

## Arguments

... One or more vector-like or matrix-like objects.

na.rm See `?base::pmax` for a description of this argument.

## Value

See `?base::pmax` for the value returned by the default methods.

Specific methods defined in Bioconductor packages will typically return an object of the same class as the input objects.

## See Also

- `base::pmax` for the default `pmax`, `pmin`, `pmax.int` and `pmin.int` methods.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `pmax,Rle-method` in the **S4Vectors** package for an example of a specific `pmax` method (defined for `Rle` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
pmax
showMethods("pmax")
selectMethod("pmax", "ANY") # the default method

pmin
showMethods("pmin")
selectMethod("pmin", "ANY") # the default method

pmax.int
showMethods("pmax.int")
selectMethod("pmax.int", "ANY") # the default method

pmin.int
showMethods("pmin.int")
selectMethod("pmin.int", "ANY") # the default method
```

---

fileName	<i>Accessing the file name of an object</i>
----------	---

---

### Description

Get the file name of an object.

### Usage

```
fileName(object, ...)
```

### Arguments

object	An object with a file name.
...	Additional arguments, for use in specific methods.

### See Also

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [fileName,MSmap-method](#) in the **MSnbase** package for an example of a specific fileName method (defined for **MSmap** objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

### Examples

```
fileName
showMethods("fileName")

library(MSnbase)
showMethods("fileName")
selectMethod("fileName", "MSmap")
```

---

format	<i>Format an R object for pretty printing</i>
--------	---

---

### Description

Turn an R object into a character vector used for pretty printing.

NOTE: This man page is for the *format S4 generic function* defined in the **BiocGenerics** package. See `?base::format` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
format(x, ...)
```

## Arguments

x	The object to format.
...	Additional arguments, for use in specific methods.

## Value

A character vector that provides a "compact representation" of x. This character vector is typically used by `print.data.frame` to display the columns of a `data.frame` object. See `?base::print.data.frame` for more information.

## See Also

- `base::format` for the default format method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `BiocGenerics` for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
format
showMethods("format")
selectMethod("format", "ANY") # the default method
```

## Description

Reduce uses a binary function to successively combine the elements of a given list-like or vector-like object and a possibly given initial value. Filter extracts the elements of a list-like or vector-like object for which a predicate (logical) function gives true. Find and Position give the first or last such element and its position in the object, respectively. Map applies a function to the corresponding elements of given list-like or vector-like objects.

NOTE: This man page is for the Reduce, Filter, Find, Map and Position *S4 generic functions* defined in the **BiocGenerics** package. See `?base::Reduce` for the default methods (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically list-like or vector-like) not supported by the default methods.

**Usage**

```
Reduce(f, x, init, right=FALSE, accumulate=FALSE, simplify=TRUE)
Filter(f, x)
Find(f, x, right=FALSE, nomatch=NULL)
Map(f, ...)
Position(f, x, right=FALSE, nomatch=NA_integer_)
```

**Arguments**

```
f, init, right, accumulate, nomatch, simplify
    See ?base::Reduce for a description of these arguments.
x
    A list-like or vector-like object.
...
    One or more list-like or vector-like objects.
```

**Value**

See ?base::Reduce for the value returned by the default methods.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default methods.

**See Also**

- base::Reduce for the default Reduce, Filter, Find, Map and Position methods.
- showMethods for displaying a summary of the methods defined for a given generic function.
- selectMethod for getting the definition of a specific method.
- Reduce,List-method in the **S4Vectors** package for an example of a specific Reduce method (defined for List objects).
- BiocGenerics for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
Reduce # note the dispatch on the 'x' arg only
showMethods("Reduce")
selectMethod("Reduce", "ANY") # the default method

Filter # note the dispatch on the 'x' arg only
showMethods("Filter")
selectMethod("Filter", "ANY") # the default method

Find # note the dispatch on the 'x' arg only
showMethods("Find")
selectMethod("Find", "ANY") # the default method

Map # note the dispatch on the '...' arg only
showMethods("Map")
selectMethod("Map", "ANY") # the default method

Position # note the dispatch on the 'x' arg only
```

```
showMethods("Position")
selectMethod("Position", "ANY") # the default method
```

---

get *Return the value of a named object*

---

## Description

Search for an object with a given name and return it.

NOTE: This man page is for the `get` and `mget` *S4 generic functions* defined in the **BiocGenerics** package. See `?base::get` for the default methods (defined in the **base** package). Bioconductor packages can define specific methods for objects (list-like or environment-like) not supported by the default methods.

## Usage

```
get(x, pos=-1, envir=as.environment(pos), mode="any", inherits=TRUE)
mget(x, envir, mode="any", ifnotfound, inherits=FALSE)
```

## Arguments

x	For <code>get</code> : A variable name (or, more generally speaking, a <i>key</i> ), given as a single string. For <code>mget</code> : A vector of variable names (or <i>keys</i> ).
envir	Where to look for the key(s). Typically a list-like or environment-like object.
pos, mode, inherits, ifnotfound	See <code>?base::get</code> for a description of these arguments.

## Details

See `?base::get` for details about the default methods.

## Value

For `get`: The value corresponding to the specified key.

For `mget`: The list of values corresponding to the specified keys. The returned list must have one element per key, and in the same order as in `x`.

See `?base::get` for the value returned by the default methods.

## See Also

- `base::get` for the default `get` and `mget` methods.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `get,ANY,Bimap,missing-method` in the **AnnotationDbi** package for an example of a specific `get` method (defined for **Bimap** objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
get # note the dispatch on the 'x', 'pos' and 'envir' args only
showMethods("get")
selectMethod("get", c("ANY", "ANY", "ANY")) # the default method

mget # note the dispatch on the 'x' and 'envir' args only
showMethods("mget")
selectMethod("mget", c("ANY", "ANY")) # the default method
```

---

grep

*Pattern Matching and Replacement*

---

## Description

Search for matches to argument 'pattern' within each element of a character vector.

NOTE: This man page is for the `grep` and `grep1` *S4 generic functions* defined in the **BiocGenerics** package. See `?base::grep` for the default methods (defined in the **base** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
grep(pattern, x, ignore.case = FALSE, perl = FALSE, value = FALSE,
      fixed = FALSE, useBytes = FALSE, invert = FALSE)
grep1(pattern, x, ignore.case = FALSE, perl = FALSE,
       fixed = FALSE, useBytes = FALSE)
```

## Arguments

pattern	The pattern for searching in x, such as a regular expression.
x	The character vector (in the general sense) to search.
ignore.case, perl, value, fixed, useBytes, invert	See <code>?base::grep</code> for a description of these arguments.

## Value

See `?base::grep` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## See Also

- `base::grep` for the default `grep` and `grep1` methods.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
grep # note the dispatch on 'pattern' and 'x' args only
showMethods("grep")
selectMethod("grep", "ANY") # the default method
```

---

image	<i>Display a color image</i>
-------	------------------------------

---

## Description

Creates a grid of colored or gray-scale rectangles with colors corresponding to the values in *z*. This can be used to display three-dimensional or spatial data aka *images*.

NOTE: This man page is for the *image S4 generic function* defined in the **BiocGenerics** package. See `?graphics::image` for the default method (defined in the **graphics** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
image(x, ...)
```

## Arguments

*x*, ... See `?graphics::image`.

## Details

See `?graphics::image` for the details.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## See Also

- `graphics::image` for the default image method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `image,AffyBatch-method` in the **affy** package for an example of a specific image method (defined for `AffyBatch` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
image
showMethods("image")
selectMethod("image", "ANY") # the default method

library(affy)
showMethods("image")
## The image() method for AffyBatch objects:
selectMethod("image", "AffyBatch")
```

---

IQR

*The Interquartile Range*

---

## Description

Compute the interquartile range for a vector.

NOTE: This man page is for the IQR *S4 generic function* defined in the **BiocGenerics** package. See `?stats::IQR` for the default method (defined in the **stats** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
IQR(x, na.rm = FALSE, type = 7)
```

## Arguments

`x`, `na.rm`, `type` See `?stats::IQR`.

## Value

See `?stats::IQR` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## See Also

- `stats::IQR` for the default IQR method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
IQR
showMethods("IQR")
selectMethod("IQR", "ANY") # the default method
```

---

`is.unsorted`*Test if a vector-like object is not sorted*

---

## Description

Test if a vector-like object is not sorted, without the cost of sorting it.

NOTE: This man page is for the `is.unsorted` *S4 generic function* defined in the **BiocGenerics** package. See `?base::is.unsorted` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-like) not supported by the default method.

## Usage

```
is.unsorted(x, na.rm=FALSE, strictly=FALSE, ...)
```

## Arguments

`x` A vector-like object.

`na.rm, strictly` See `?base::is.unsorted` for a description of these arguments.

`...` Additional arguments, for use in specific methods.

Note that `base::is.unsorted` (the default method) only takes the `x`, `na.rm`, and `strictly` arguments.

## Value

See `?base::is.unsorted` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## Note

TO DEVELOPERS:

The `is.unsorted` method for specific vector-like objects should adhere to the same underlying order used by the `order`, `sort`, and `rank` methods for the same objects.

## See Also

- `base::is.unsorted` for the default `is.unsorted` method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `is.unsorted,GenomicRanges-method` in the **GenomicRanges** package for an example of a specific `is.unsorted` method (defined for `GenomicRanges` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
is.unsorted # note the dispatch on the 'x' arg only
showMethods("is.unsorted")
selectMethod("is.unsorted", "ANY") # the default method
```

---

lapply

*Apply a function over a list-like or vector-like object*


---

**Description**

lapply returns a list of the same length as X, each element of which is the result of applying FUN to the corresponding element of X.

sapply is a user-friendly version and wrapper of lapply by default returning a vector, matrix or, if simplify="array", an array if appropriate, by applying simplify2array(). sapply(x, f, simplify=FALSE, USE.NAMES=FALSE) is the same as lapply(x, f).

NOTE: This man page is for the lapply and sapply *S4 generic functions* defined in the **BiocGenerics** package. See `?base::lapply` for the default methods (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically list-like or vector-like) not supported by the default methods.

**Usage**

```
lapply(X, FUN, ...)
sapply(X, FUN, ..., simplify=TRUE, USE.NAMES=TRUE)
```

**Arguments**

X                           A list-like or vector-like object.  
 FUN, ..., simplify, USE.NAMES  
                           See `?base::lapply` for a description of these arguments.

**Value**

See `?base::lapply` for the value returned by the default methods.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default methods. In particular, lapply and sapply(simplify=FALSE) should always return a list.

**See Also**

- `base::lapply` for the default lapply and sapply methods.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `lapply,List-method` in the **S4Vectors** package for an example of a specific lapply method (defined for `List` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
lapply # note the dispatch on the 'X' arg only
showMethods("lapply")
selectMethod("lapply", "ANY") # the default method
```

```
sapply # note the dispatch on the 'X' arg only
showMethods("sapply")
selectMethod("sapply", "ANY") # the default method
```

---

mad

*Median Absolute Deviation*

---

## Description

Compute the median absolute deviation for a vector, dispatching only on the first argument, *x*.

NOTE: This man page is for the *mad S4 generic function* defined in the **BiocGenerics** package. See `?stats::mad` for the default method (defined in the **stats** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
mad(x, center = median(x), constant = 1.4826,
    na.rm = FALSE, low = FALSE, high = FALSE)
```

## Arguments

*x*, *center*, *constant*, *na.rm*, *low*, *high*  
See `?stats::mad`.

## Value

See `?stats::mad` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## See Also

- `stats::mad` for the default *mad* method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
mad
showMethods("mad")
selectMethod("mad", "ANY") # the default method
```

---

`mapply`*Apply a function to multiple list-like or vector-like arguments*

---

## Description

`mapply` is a multivariate version of `sapply`. `mapply` applies `FUN` to the first elements of each ... argument, the second elements, the third elements, and so on. Arguments are recycled if necessary.

NOTE: This man page is for the `mapply` *S4 generic function* defined in the **BiocGenerics** package. See `?base::mapply` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically list-like or vector-like) not supported by the default methods.

## Usage

```
mapply(FUN, ..., MoreArgs=NULL, SIMPLIFY=TRUE, USE.NAMES=TRUE)
```

## Arguments

`FUN`, `MoreArgs`, `SIMPLIFY`, `USE.NAMES`

See `?base::mapply` for a description of these arguments.

... One or more list-like or vector-like objects of strictly positive length, or all of zero length.

## Value

See `?base::mapply` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## See Also

- `base::mapply` for the default `mapply` method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
mapply # note the dispatch on the '...' arg only
showMethods("mapply")
selectMethod("mapply", "ANY") # the default method
```

---

match	Value matching
-------	----------------

---

## Description

`match` returns a vector of the positions of (first) matches of its first argument in its second.

`%in%` is a binary operator that returns a logical vector of the length of its left operand indicating if the elements in it have a match or not.

NOTE: This man page is for the `match` and `%in%` *S4 generic functions* defined in the **BiocGenerics** package. See `?base::match` for the default methods (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-like) not supported by the default methods.

## Usage

```
match(x, table, nomatch=NA_integer_, incomparables=NULL, ...)
```

```
x %in% table
```

## Arguments

`x, table` Vector-like objects (typically of the same class, but not necessarily).

`nomatch, incomparables`

See `?base::match` for a description of these arguments.

`...` Additional arguments, for use in specific methods.

## Value

The same as the default methods (see `?base::match` for the value returned by the default methods).

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default methods.

## Note

The default `base::match` method (defined in the **base** package) doesn't have the `...` argument. We've added it to the generic function defined in the **BiocGenerics** package in order to allow specific methods to support additional arguments if needed.

## See Also

- `base::match` for the default `match` and `%in%` methods.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `match,Hits,Hits-method` and `%in%,Rle,ANY-method` in the **S4Vectors** package for examples of specific `match` and `%in%` methods (defined for `Hits` and `Rle` objects, respectively).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```

match # note the dispatch on the 'x' and 'table' args only
showMethods("match")
selectMethod("match", c("ANY", "ANY")) # the default method

`%in%`
showMethods("%in%")
selectMethod("%in%", c("ANY", "ANY")) # the default method

```

---

mean

*Arithmetic Mean*

---

**Description**

Generic function for the (trimmed) arithmetic mean.

NOTE: This man page is for the mean *S4 generic function* defined in the **BiocGenerics** package. See `?base::mean` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-like) not supported by the default method.

**Usage**

```
mean(x, ...)
```

**Arguments**

x typically a vector-like object  
 ... see [mean](#)

**Value**

See `?base::mean` for the value returned by the default method.

Specific methods defined in Bioconductor packages will typically return an object of the same class as the input object.

**See Also**

- `base::mean` for the default mean method.
- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [mean,Rle-method](#) in the **S4Vectors** package for an example of a specific mean method (defined for **Rle** objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```

mean
showMethods("mean")
selectMethod("mean", "ANY") # the default method

```

---

normalize	<i>Normalize an object</i>
-----------	----------------------------

---

### Description

A generic function which normalizes an object containing microarray data or other data. Normalization is intended to remove from the intensity measures any systematic trends which arise from the microarray technology rather than from differences between the probes or between the target RNA samples hybridized to the arrays.

### Usage

```
normalize(object, ...)
```

### Arguments

object	A data object, typically containing microarray data.
...	Additional arguments, for use in specific methods.

### Value

An object containing the normalized data.

### See Also

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [normalize,AffyBatch-method](#) in the **affy** package and [normalize,MSnExp-method](#) in the **MSnbase** package for examples of specific normalize methods (defined for [AffyBatch](#) and [MSnExp](#) objects, respectively).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

### Examples

```
normalize
showMethods("normalize")

library(affy)
showMethods("normalize")
selectMethod("normalize", "AffyBatch")
```

---

nrow	<i>The number of rows/columns of an array-like object</i>
------	---

---

## Description

Return the number of rows or columns present in an array-like object.

NOTE: This man page is for the `nrow`, `ncol`, `NROW` and `NCOL` *S4 generic functions* defined in the **BiocGenerics** package. See `?base::nrow` for the default methods (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically matrix- or array-like) not supported by the default methods.

## Usage

```
nrow(x)
ncol(x)
NROW(x)
NCOL(x)
```

## Arguments

`x`                    A matrix- or array-like object.

## Value

A single integer or NULL.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default methods.

## See Also

- `base::nrow` for the default `nrow`, `ncol`, `NROW` and `NCOL` methods.
- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [nrow,DataFrame-method](#) in the **S4Vectors** package for an example of a specific `nrow` method (defined for [DataFrame](#) objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
nrow
showMethods("nrow")
selectMethod("nrow", "ANY") # the default method

ncol
showMethods("ncol")
selectMethod("ncol", "ANY") # the default method
```

```
NROW
showMethods("NROW")
selectMethod("NROW", "ANY") # the default method

NCOL
showMethods("NCOL")
selectMethod("NCOL", "ANY") # the default method
```

---

Ontology	<i>Generic Ontology getter</i>
----------	--------------------------------

---

### Description

Get the Ontology of an object.

### Usage

```
Ontology(object)
```

### Arguments

object            An object with an Ontology.

### See Also

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [Ontology, GOTerms-method](#) in the **AnnotationDbi** package for an example of a specific Ontology method (defined for [GOTerms](#) objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

### Examples

```
Ontology
showMethods("Ontology")

library(AnnotationDbi)
showMethods("Ontology")
selectMethod("Ontology", "GOTerms")
```

---

order	<i>Ordering permutation</i>
-------	-----------------------------

---

## Description

`order` returns a permutation which rearranges its first argument into ascending or descending order, breaking ties by further arguments.

NOTE: This man page is for the `order` *S4 generic function* defined in the **BiocGenerics** package. See `?base::order` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-like) not supported by the default method.

## Usage

```
order(..., na.last=TRUE, decreasing=FALSE, method=c("auto", "shell", "radix"))
```

## Arguments

`...` One or more vector-like objects, all of the same length.  
`na.last`, `decreasing`, `method`  
See `?base::order` for a description of these arguments.

## Value

The default method (see `?base::order`) returns an integer vector of length `N` where `N` is the common length of the input objects. This integer vector represents a permutation of `N` elements and can be used to rearrange the first argument in `...` into ascending or descending order (by subsetting it).

Specific methods defined in Bioconductor packages should also return an integer vector representing a permutation of `N` elements.

## Note

TO DEVELOPERS:

Specific `order` methods should preferably be made "stable" for consistent behavior across platforms and consistency with `base::order()`. Note that C `qsort()` is *not* "stable" so `order` methods that use `qsort()` at the C-level need to ultimately break ties by position, which can easily be done by adding a little extra code at the end of the comparison function passed to `qsort()`.

`order(x, decreasing=TRUE)` is *not* always equivalent to `rev(order(x))`.

`order`, `sort`, and `rank` methods for specific vector-like objects should adhere to the same underlying order that should be conceptually defined as a binary relation on the set of all possible vector values. For completeness, this binary relation should also be incarnated by a `<=` method.

### See Also

- `base::order` for the default order method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `order,IntegerRanges-method` in the **IRanges** package for an example of a specific order method (defined for `IntegerRanges` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

### Examples

```
order
showMethods("order")
selectMethod("order", "ANY") # the default method
```

---

organism_species	<i>Organism and species accessors</i>
------------------	---------------------------------------

---

### Description

Get or set the organism and/or species of an object.

### Usage

```
organism(object)
organism(object) <- value

species(object)
species(object) <- value
```

### Arguments

object	An object to get or set the organism or species of.
value	The organism or species to set on object.

### Value

`organism` should return the *scientific name* (i.e. genus and species, or genus and species and sub-species) of the organism. Preferably in the format "Genus species" (e.g. "Homo sapiens") or "Genus species subspecies" (e.g. "Homo sapiens neanderthalensis").

`species` should of course return the species of the organism. Unfortunately there is a long history of misuse of this accessor in Bioconductor so its usage is now discouraged (starting with BioC 3.1).

## Note

### TO DEVELOPERS:

species has been historically misused in many places in Bioconductor and is redundant with organism. So implementing the species accessor is now discouraged (starting with BioC 3.1). The organism accessor (returning the *scientific name*) should be implemented instead.

## See Also

- [http://bioconductor.org/packages/release/BiocViews.html#\\_\\_\\_Organism](http://bioconductor.org/packages/release/BiocViews.html#___Organism) for browsing the annotation packages currently available in Bioconductor by organism.
- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [organism,character-method](#) and [organism,chromLocation-method](#) in the **annotate** package for examples of specific organism methods (defined for [character](#) and [chromLocation](#) objects).
- [species,AnnotationDb-method](#) in the **AnnotationDbi** package for an example of a specific species method (defined for [AnnotationDb](#) objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
## organism() getter:
organism
showMethods("organism")

library(annotate)
showMethods("organism")
selectMethod("organism", "character")
selectMethod("organism", "chromLocation")

## organism() setter:
`organism<-`
showMethods("organism<-")

## species() getter:
species
showMethods("species")

library(AnnotationDbi)
selectMethod("species", "AnnotationDb")

## species() setter:
`species<-`
showMethods("species<-")
```

---

paste	<i>Concatenate strings</i>
-------	----------------------------

---

## Description

paste concatenates vectors of strings or vector-like objects containing strings.

NOTE: This man page is for the `paste S4 generic function` defined in the **BiocGenerics** package. See `?base::paste` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-like objects containing strings) not supported by the default method.

## Usage

```
paste(..., sep=" ", collapse=NULL, recycle0=FALSE)
```

## Arguments

... One or more vector-like objects containing strings.  
sep, collapse, recycle0  
See `?base::paste` for a description of these arguments.

## Value

See `?base::paste` for the value returned by the default method.

Specific methods defined in Bioconductor packages will typically return an object of the same class as the input objects.

## See Also

- `base::paste` for the default paste method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `paste,Rle-method` in the **S4Vectors** package for an example of a specific paste method (defined for **Rle** objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
paste
showMethods("paste")
selectMethod("paste", "ANY") # the default method
```

---

path *Accessing the path of an object*

---

**Description**

Get or set the path of an object.

**Usage**

```
path(object, ...)  
path(object, ...) <- value
```

```
basename(path, ...)  
basename(path, ...) <- value
```

```
dirname(path, ...)  
dirname(path, ...) <- value
```

```
## The purpose of the following methods is to make the basename() and  
## dirname() getters work out-of-the-box on any object for which the  
## path() getter works.
```

```
## S4 method for signature 'ANY'  
basename(path, ...)
```

```
## S4 method for signature 'ANY'  
dirname(path, ...)
```

```
## The purpose of the following replacement methods is to make the  
## basename() and dirname() setters work out-of-the-box on any object  
## for which the path() getter and setter work.
```

```
## S4 replacement method for signature 'character'  
basename(path, ...) <- value
```

```
## S4 replacement method for signature 'ANY'  
basename(path, ...) <- value
```

```
## S4 replacement method for signature 'character'  
dirname(path, ...) <- value
```

```
## S4 replacement method for signature 'ANY'  
dirname(path, ...) <- value
```

**Arguments**

object            An object containing paths. Even though it will typically contain a single path, object can actually contain an arbitrary number of paths.

...	Additional arguments, for use in specific methods.
value	For path<- , the paths to set on object. For basename<- or dirname<- , the basenames or dirnames to set on path.
path	A character vector <i>or an object containing paths</i> .

### Value

A character vector for path(object), basename(path), and dirname(path). Typically of length 1 but not necessarily. Possibly with names on it for path(object).

### See Also

- base::[basename](#) for the functions the basename and dirname generics are based on.
- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [path,RsamtoolsFile-method](#) in the **Rsamtools** package for an example of a specific path method (defined for [RsamtoolsFile](#) objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

### Examples

```
## -----
## GENERIC FUNCTIONS AND DEFAULT METHODS
## -----

path
showMethods("path")

`path<-`
showMethods("path<-")

basename
showMethods("basename")

`basename<-`
showMethods("basename<-")

dirname
showMethods("dirname")

`dirname`
showMethods("dirname<-")

## Default basename() and dirname() getters:
selectMethod("basename", "ANY")
selectMethod("dirname", "ANY")

## Default basename() and dirname() setters:
selectMethod("basename<-", "character")
```

```

selectMethod("basename<-", "ANY")
selectMethod("dirname<-", "character")
selectMethod("dirname<-", "ANY")

## -----
## OBJECTS CONTAINING PATHS
## -----

## Let's define a simple class to represent objects that contain paths:
setClass("A", slots=c(stuff="ANY", path="character"))

a <- new("A", stuff=runif(5),
         path=c(one="path/to/file1", two="path/to/file2"))

## path() getter:
setMethod("path", "A", function(object) object@path)

path(a)

## Because the path() getter works on 'a', now the basename() and
## dirname() getters also work:
basename(a)
dirname(a)

## path() setter:
setReplaceMethod("path", "A",
  function(object, ..., value)
  {
    if (length(list(...)) != 0L) {
      dots <- match.call(expand.dots=FALSE)[[3L]]
      stop(BiocGenerics:::unused_arguments_msg(dots))
    }
    object@path <- value
    object
  }
)

a <- new("A", stuff=runif(5))
path(a) <- c(one="path/to/file1", two="path/to/file2")
path(a)

## Because the path() getter and setter work on 'a', now the basename()
## and dirname() setters also work:
basename(a) <- toupper(basename(a))
path(a)
dirname(a) <- "~/MyDataFiles"
path(a)

```

**Description**

A generic function which produces an MA-plot for an object containing microarray, RNA-Seq or other data.

**Usage**

```
plotMA(object, ...)
```

**Arguments**

object	A data object, typically containing count values from an RNA-Seq experiment or microarray intensity values.
...	Additional arguments, for use in specific methods.

**Value**

Undefined. The function exists for its side effect, producing a plot.

**See Also**

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [plotMA](#) in the **limma** package for a function with the same name that is not dispatched through this generic function.
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
showMethods("plotMA")

suppressWarnings(
  if(require("DESeq2"))
    example("plotMA", package="DESeq2", local=TRUE)
)
```

---

plotPCA

*PCA-plot: Principal Component Analysis plot*

---

**Description**

A generic function which produces a PCA-plot.

**Usage**

```
plotPCA(object, ...)
```



## Arguments

- `x` A vector-like object.
- `na.last`, `ties.method` See `?base::rank` for a description of these arguments.
- `...` Additional arguments, for use in specific methods.
- Note that `base::rank` (the default method) only takes the `x`, `na.last`, and `ties.method` arguments.

## Value

See `?base::rank` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## Note

TO DEVELOPERS:

See note in `?BiocGenerics::order` about "stable" order.

`order`, `sort`, and `rank` methods for specific vector-like objects should adhere to the same underlying order that should be conceptually defined as a binary relation on the set of all possible vector values. For completeness, this binary relation should also be incarnated by a `<=` method.

## See Also

- `base::rank` for the default rank method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `rank, Vector-method` in the **S4Vectors** package for an example of a specific rank method (defined for `Vector` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
rank # note the dispatch on the 'x' arg only
showMethods("rank")
selectMethod("rank", "ANY") # the default method
```

---

`relist`*Re-listing an unlist()ed object*

---

## Description

`relist` is a generic function with a few methods in order to allow easy inversion of `unlist(x)`.

NOTE: This man page is for the `relist` *S4 generic function* defined in the **BiocGenerics** package. See `?utils::relist` for the default method (defined in the **utils** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
relist(flesh, skeleton)
```

## Arguments

<code>flesh</code>	A vector-like object.
<code>skeleton</code>	A list-like object. Only the "shape" (i.e. the lengths of the individual list elements) of <code>skeleton</code> matters. Its exact content is ignored.

## Value

A list-like object with the same "shape" as `skeleton` and that would give `flesh` back if `unlist()`d.

## See Also

- `utils::relist` for the default `relist` method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `relist,ANY,List-method` in the **IRanges** package for an example of a specific `relist` method (defined for when `skeleton` is a `List` object).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
relist
showMethods("relist")
selectMethod("relist", c("ANY", "ANY")) # the default method
```

---

`rep`*Replicate elements of a vector-like object*

---

## Description

`rep.int` replicates the elements in `x`.

NOTE: This man page is for the `rep.int` *S4 generic function* defined in the **BiocGenerics** package. See `?base::rep.int` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-like) not supported by the default method.

## Usage

```
rep.int(x, times)
```

## Arguments

<code>x</code>	The object to replicate (typically vector-like).
<code>times</code>	See <code>?base::rep.int</code> for a description of this argument.

## Value

See `?base::rep.int` for the value returned by the default method.

Specific methods defined in Bioconductor packages will typically return an object of the same class as the input object.

## See Also

- `base::rep.int` for the default `rep.int`, `intersect`, and `setdiff` methods.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `rep.int,Rle-method` in the **S4Vectors** package for an example of a specific `rep.int` method (defined for **Rle** objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
rep.int
showMethods("rep.int")
selectMethod("rep.int", "ANY") # the default method
```

---

residuals	<i>Extract model residuals</i>
-----------	--------------------------------

---

### Description

`residuals` is a generic function which extracts model residuals from objects returned by modeling functions.

NOTE: This man page is for the `residuals` *S4 generic function* defined in the **BiocGenerics** package. See `?stats::residuals` for the default method (defined in the **stats** package). Bioconductor packages can define specific methods for objects not supported by the default method.

### Usage

```
residuals(object, ...)
```

### Arguments

`object, ...` See `?stats::residuals`.

### Value

Residuals extracted from the object `object`.

### See Also

- `stats::residuals` for the default residuals method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `residuals,PLMset-method` in the **affyPLM** package for an example of a specific residuals method (defined for `PLMset` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

### Examples

```
residuals
showMethods("residuals")
selectMethod("residuals", "ANY") # the default method
```

---

row+colnames	<i>Row and column names</i>
--------------	-----------------------------

---

## Description

Get or set the row or column names of a matrix-like object.

NOTE: This man page is for the `rownames`, ``rownames<-``, `colnames`, and ``colnames<-`` *S4 generic functions* defined in the **BiocGenerics** package. See `?base::rownames` for the default methods (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically matrix-like) not supported by the default methods.

## Usage

```
rownames(x, do.NULL=TRUE, prefix="row")
rownames(x) <- value
```

```
colnames(x, do.NULL=TRUE, prefix="col")
colnames(x) <- value
```

## Arguments

<code>x</code>	A matrix-like object.
<code>do.NULL</code> , <code>prefix</code>	See <code>?base::rownames</code> for a description of these arguments.
<code>value</code>	Either NULL or a character vector equal of length equal to the appropriate dimension.

## Value

The getters will return NULL or a character vector of length `nrow(x)` for `rownames` and length `ncol(x)` for `colnames(x)`.

See `?base::rownames` for more information about the default methods, including how the setters are expected to behave.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default methods.

## See Also

- `base::rownames` for the default `rownames`, ``rownames<-``, `colnames`, and ``colnames<-`` methods.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `rownames,DataFrame-method` in the **S4Vectors** package for an example of a specific `rownames` method (defined for `DataFrame` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
## rownames() getter:
rownames # note the dispatch on the 'x' arg only
showMethods("rownames")
selectMethod("rownames", "ANY") # the default method

## rownames() setter:
`rownames<-`
showMethods("rownames<-")
selectMethod("rownames<-", "ANY") # the default method

## colnames() getter:
colnames # note the dispatch on the 'x' arg only
showMethods("colnames")
selectMethod("colnames", "ANY") # the default method

## colnames() setter:
`colnames<-`
showMethods("colnames<-")
selectMethod("colnames<-", "ANY") # the default method
```

---

S3-classes-as-S4-classes

*S3 classes as S4 classes*

---

## Description

Some old-style (aka S3) classes are turned into formally defined (aka S4) classes by the **Bioc-Generics** package. This allows S4 methods defined in Bioconductor packages to use them in their signatures.

## Details

S3 classes currently turned into S4 classes:

- connection class and subclasses: [connection](#), file, url, gzfile, bzfile, unz, pipe, fifo, sockconn, terminal, textConnection, gzcon. Additionally the `character_OR_connection` S4 class is defined as the union of classes `character` and `connection`.
- others: [AsIs](#), [dist](#)

## See Also

[setOldClass](#) and [setClassUnion](#) in the **methods** package.

---

saveRDS

*The saveRDS() S4 generic and default method*

---

## Description

Generic function to write a single R object to a file.

NOTE: This man page is for the `saveRDS` S4 generic function and default method defined in the **BiocGenerics** package. See `?base::saveRDS` for the corresponding function defined in base R.

## Usage

```
saveRDS(object, file="", ascii=FALSE, version=NULL,
        compress=TRUE, rehook=NULL)
```

## Arguments

`object`, `file`, `ascii`, `version`, `compress`, `rehook`

See `?base::saveRDS` for a description of these arguments.

## Details

The default `saveRDS` method defined in this package is a thin wrapper around `base::saveRDS` that issues a warning if the object to serialize contains out-of-memory data. See `?containsOutOfMemoryData` for more information.

Bioconductor packages can override this default method with more specialized methods.

## Value

An invisible NULL.

## See Also

- `base::saveRDS` in the **base** package for the default `saveRDS` method.
- `containsOutOfMemoryData` for determining whether an object contains out-of-memory data or not.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `saveRDS, SummarizedExperiment-method` in the **SummarizedExperiment** package for an example of a specific `saveRDS` method (defined for `SummarizedExperiment` objects and derivatives).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
saveRDS # note the dispatch on the 'object' arg only
showMethods("saveRDS")
selectMethod("saveRDS", "ANY") # the default method
```

---

score	<i>Score accessor</i>
-------	-----------------------

---

## Description

Get or set the score value contained in an object.

## Usage

```
score(x, ...)  
score(x, ...) <- value
```

## Arguments

x	An object to get or set the score value of.
...	Additional arguments, for use in specific methods.
value	The score value to set on x.

## See Also

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [score.GenomicRanges-method](#) in the **GenomicRanges** package for an example of a specific score method (defined for [GenomicRanges](#) objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
score  
showMethods("score")  
  
`score<-`  
showMethods("score<-")  
  
library(GenomicRanges)  
  
showMethods("score")  
selectMethod("score", "GenomicRanges")  
  
showMethods("score<-")  
selectMethod("score<-", "GenomicRanges")
```

## Description

Performs *set* union, intersection and (asymmetric!) difference on two vector-like objects.

NOTE: This man page is for the `union`, `intersect` and `setdiff` *S4 generic functions* defined in the **BiocGenerics** package. See `?base::union` for the default methods (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-like) not supported by the default methods.

## Usage

```
union(x, y, ...)  
intersect(x, y, ...)  
setdiff(x, y, ...)
```

## Arguments

<code>x, y</code>	Vector-like objects (typically of the same class, but not necessarily).
<code>...</code>	Additional arguments, for use in specific methods.

## Value

See `?base::union` for the value returned by the default methods.

Specific methods defined in Bioconductor packages will typically return an object of the same class as the input objects.

## Note

The default methods (defined in the **base** package) only take 2 arguments. We've added the `...` argument to the generic functions defined in the **BiocGenerics** package so they can be called with an arbitrary number of effective arguments. For `union` or `intersect`, this typically allows Bioconductor packages to define methods that compute the union or intersection of more than 2 objects. However, for `setdiff`, which is conceptually a binary operation, this typically allows methods to add extra arguments for controlling/altering the behavior of the operation. Like for example the `ignore.strand` argument supported by the `setdiff` method for **GenomicRanges** objects (defined in the **GenomicRanges** package). (Note that the `union` and `intersect` methods for those objects also support the `ignore.strand` argument.)

## See Also

- `base::union` for the default `union`, `intersect`, and `setdiff` methods.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.

- [union, GenomicRanges, GenomicRanges-method](#) in the **GenomicRanges** package for examples of specific union, intersect, and setdiff methods (defined for [GenomicRanges](#) objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
union
showMethods("union")
selectMethod("union", c("ANY", "ANY")) # the default method

intersect
showMethods("intersect")
selectMethod("intersect", c("ANY", "ANY")) # the default method

setdiff
showMethods("setdiff")
selectMethod("setdiff", c("ANY", "ANY")) # the default method
```

---

sort

*Sorting a vector-like object*

---

## Description

Sort a vector-like object into ascending or descending order.

NOTE: This man page is for the *sort S4 generic function* defined in the **BiocGenerics** package. See `?base::sort` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
sort(x, decreasing=FALSE, ...)
```

## Arguments

`x` A vector-like object.

`decreasing, ...` See `?base::sort` for a description of these arguments.

## Value

See `?base::sort` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

**Note**

TO DEVELOPERS:

See note in `?BiocGenerics::order` about "stable" order.

`order`, `sort`, and `rank` methods for specific vector-like objects should adhere to the same underlying order that should be conceptually defined as a binary relation on the set of all possible vector values. For completeness, this binary relation should also be incarnated by a `<=` method.

**See Also**

- `base::sort` for the default sort method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `sort,Vector-method` in the **S4Vectors** package for an example of a specific sort method (defined for `Vector` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
sort # note the dispatch on the 'x' arg only
showMethods("sort")
selectMethod("sort", "ANY") # the default method
```

---

start

*The start(), end(), width(), and pos() generic getters and setters*

---

**Description**

Get or set the start, end, width, or single positions stored in an object.

NOTE: This man page is for the `start`, ``start<-``, `end`, ``end<-``, `width`, ``width<-``, and `pos` *S4 generic functions* defined in the **BiocGenerics** package. See `?stats::start` for the `start` and `end` *S3 generics* defined in the **stats** package.

**Usage**

```
start(x, ...)
start(x, ...) <- value

end(x, ...)
end(x, ...) <- value

width(x)
width(x, ...) <- value

pos(x)
```

## Arguments

x	For the <code>start()</code> , <code>end()</code> , and <code>width()</code> getters/setters: an object containing start, end, and width values. For the <code>pos{}</code> getter: an object containing single positions.
...	Additional arguments, for use in specific methods.
value	The start, end, or width values to set on x.

## Value

See specific methods defined in Bioconductor packages.

## See Also

- `stats::start` in the **stats** package for the start and end S3 generics.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `start,IRanges-method` in the **IRanges** package for examples of specific start, end, and width methods (defined for **IRanges** objects).
- `pos,UnstitchedIPos-method` in the **IRanges** package for an example of a specific pos method (defined for **UnstitchedIPos** objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
## start() getter:
start
showMethods("start")

library(IRanges)
showMethods("start")
selectMethod("start", "IRanges") # start() getter for IRanges objects

## start() setter:
`start<-`
showMethods("start<-")
selectMethod("start<-", "IRanges") # start() setter for IRanges objects

## end() getter:
end
showMethods("end")
selectMethod("end", "IRanges") # end() getter for IRanges objects

## end() setter:
`end<-`
showMethods("end<-")
selectMethod("end<-", "IRanges") # end() setter for IRanges objects

## width() getter:
```

```

width
showMethods("width")
selectMethod("width", "IRanges") # width() getter for IRanges objects

## width() setter:
`width<-`
showMethods("width<-")
selectMethod("width<-", "IRanges") # width() setter for IRanges objects

## pos() getter:
pos
showMethods("pos")
selectMethod("pos", "UnstitchedIPos") # pos() getter for UnstitchedIPos
# objects

```

---

strand	<i>Accessing strand information</i>
--------	-------------------------------------

---

## Description

Get or set the strand information contained in an object.

## Usage

```

strand(x, ...)
strand(x, ...) <- value

unstrand(x)

invertStrand(x)
## S4 method for signature 'ANY'
invertStrand(x)

```

## Arguments

x	An object containing strand information.
...	Additional arguments, for use in specific methods.
value	The strand information to set on x.

## Details

All the strand methods defined in the **GenomicRanges** package use the same set of 3 values (called the "standard strand levels") to specify the strand of a genomic location: +, -, and \*. \* is used when the exact strand of the location is unknown, or irrelevant, or when the "feature" at that location belongs to both strands.

Note that `unstrand` is not a generic function, just a convenience wrapper to the generic `strand()` setter (`strand<-`) that does:

```
strand(x) <- "*"
x
```

The default method for invertStrand does:

```
strand(x) <- invertStrand(strand(x))
x
```

### Value

If `x` is a vector-like object, `strand(x)` will typically return a vector-like object *parallel* to `x`, that is, an object of the same length as `x` where the `i`-th element describes the strand of the `i`-th element in `x`.

`unstrand(x)` and `invertStrand(x)` return a copy of `x` with the strand set to "\*" for `unstrand` or inverted for `invertStrand` (i.e. "+" and "-" switched, and "\*" untouched).

### See Also

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [strand,GRanges-method](#) in the **GenomicRanges** package for an example of a specific strand method (defined for [GRanges](#) objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

### Examples

```
strand
showMethods("strand")

`strand<-`
showMethods("strand<-")

unstrand

invertStrand
showMethods("invertStrand")
selectMethod("invertStrand", "ANY") # the default method

library(GenomicRanges)

showMethods("strand")
selectMethod("strand", "missing")
strand()

showMethods("strand<-")
```

---

subset

*Subsetting vector-like, matrix-like and data-frame-like objects*

---

## Description

Return subsets of vector-like, matrix-like or data-frame-like objects which meet conditions.

NOTE: This man page is for the subset *S4 generic function* defined in the **BiocGenerics** package.  
See `?base::subset` for the subset S3 generic defined in the **base** package.

## Usage

```
subset(x, ...)
```

## Arguments

x	A vector-like, matrix-like or data-frame-like object to be subsetted.
...	Additional arguments (e.g. subset, select, drop), for use in specific methods. See <code>?base::subset</code> for more information.

## Value

An object similar to x containing just the selected elements (for a vector-like object), or the selected rows and columns (for a matrix-like or data-frame-like object).

## See Also

- `base::subset` in the **base** package for the subset S3 generic.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `subset,RectangularData-method` in the **S4Vectors** package for an example of a specific subset method (defined for `RectangularData` derivatives).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
subset
showMethods("subset")
selectMethod("subset", "ANY") # the default method

library(S4Vectors)
showMethods("subset")
## The subset() method for RectangularData derivatives:
selectMethod("subset", "RectangularData")
```

---

`t`*Matrix Transpose*

---

**Description**

Given a rectangular object `x`, `t` returns the transpose of `x`.

NOTE: This man page is for the `t` *S4 generic function* defined in the **BiocGenerics** package. See `?base::t` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically array-like) not supported by the default method.

**Usage**

```
t(x)
```

**Arguments**

`x`                    A matrix-like or other rectangular object.

**Value**

See `?base::t` for the value returned by the default method.

Specific methods defined in Bioconductor packages will typically return an object of the same class as the input object.

**See Also**

- `base::t` for the default `t` method.
- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [t.Hits-method](#) in the **S4Vectors** package for an example of a specific `t` method (defined for [Hits](#) objects).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
t
showMethods("t")
selectMethod("t", "ANY") # the default method
```

---

table	<i>Cross tabulation and table creation</i>
-------	--

---

## Description

table uses the cross-classifying factors to build a contingency table of the counts at each combination of factor levels.

NOTE: This man page is for the table *S4 generic function* defined in the **BiocGenerics** package. See `?base::table` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
table(...)
```

## Arguments

... One or more objects which can be interpreted as factors (including character strings), or a list (or data frame) whose components can be so interpreted.

## Value

See `?base::table` for the value returned by the default method.

Specific methods defined in Bioconductor packages should also return the type of object returned by the default method.

## See Also

- `base::table` for the default table method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `table,Rle-method` in the **S4Vectors** package for an example of a specific table method (defined for **Rle** objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
table
showMethods("table")
selectMethod("table", "ANY") # the default method
```

tapply

*Apply a function over a ragged array***Description**

tapply applies a function to each cell of a ragged array, that is to each (non-empty) group of values given by a unique combination of the levels of certain factors.

NOTE: This man page is for the tapply *S4 generic function* defined in the **BiocGenerics** package. See `?base::tapply` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically list-like or vector-like) not supported by the default method.

**Usage**

```
tapply(X, INDEX, FUN=NULL, ..., default=NA, simplify=TRUE)
```

**Arguments**

X	The default method expects an atomic object, typically a vector. See <code>?base::tapply</code> for the details. Specific methods can support other objects (typically list-like or vector-like). Please refer to the documentation of a particular method for the details.
INDEX	The default method expects a list of one or more factors, each of same length as X. See <code>?base::tapply</code> for the details. Specific methods can support other objects (typically list-like). Please refer to the documentation of a particular method for the details.
FUN, ..., default, simplify	See <code>?base::tapply</code> for a description of these arguments.

**Value**

See `?base::tapply` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

**See Also**

- `base::tapply` for the default tapply method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `tapply, Vector, ANY-method` in the **IRanges** package for an example of a specific tapply method (defined for `Vector` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
tapply # note the dispatch on the 'X' and 'INDEX' args only
showMethods("tapply")
selectMethod("tapply", c("ANY", "ANY")) # the default method
```

---

testPackage	<i>Run RUnit package unit tests</i>
-------------	-------------------------------------

---

**Description**

testPackage helps developers implement unit tests using the **RUnit** testing conventions.

**Usage**

```
testPackage(pkgname=NULL, subdir="unitTests", pattern="^test_.*\\.R$",
  path=getwd())
```

**Arguments**

pkgname	The name of the package whose installed unit tests are to be run. A missing or NULL value implies that the testPackage command will look for tests within the package source directory indicated by path.
subdir	A character(1) vector providing the subdirectory in which unit tests are located. The directory is searched first in the (installed or source) package root, or in a subdirectory inst/ below the root.
pattern	A character(1) regular expression describing the file names to be evaluated; typically used to restrict tests to a subset of all test files.
path	A character(1) directory path indicating, when pkgname is missing or NULL, where unit tests will be searched. path can be any location at or below the package root.

**Details**

This function is not exported from the package namespace, and must be invoked using triple colons, `BiocGenerics:::testPackage()`; it is provided primarily for the convenience of developers.

When invoked with missing or NULL pkgname argument, the function assumes that it has been invoked from within the package source tree (or that the source tree is located above path), and finds unit tests in subdir="unitTests" in either the base or inst/ directories at the root of the package source tree. This mode is useful when developing unit tests, since the package does not have to be re-installed to run an updated test.

When invoked with pkgname set to the name of an installed package, unit tests are searched for in the installed package directory.

**Value**

The function returns the result of `RUnit::runTestSuite` invoked on the unit tests specified in the function call.

**See Also**

<http://bioconductor.org/developers/how-to/unitTesting-guidelines/>

**Examples**

```
## Run unit tests found in the library location where
## BiocGenerics is installed
BiocGenerics:::testPackage("BiocGenerics")
## Not run: ## Run unit tests for the package whose source tree implied
## by getwd()
BiocGenerics:::testPackage()

## End(Not run)
```

---

toTable	<i>An alternative to <code>as.data.frame()</code></i>
---------	---

---

**Description**

toTable() is an *S4 generic function* provided as an alternative to `as.data.frame()`.

**Usage**

```
toTable(x, ...)
```

**Arguments**

x	The object to turn into a data frame.
...	Additional arguments, for use in specific methods.

**Value**

A data frame.

**See Also**

- The `as.data.frame` *S4 generic* defined in the **BiocGenerics** package.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `toTable,Bimap-method` in the **AnnotationDbi** package for an example of a specific toTable method (defined for **Bimap** objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```

toTable
showMethods("toTable")

library(AnnotationDbi)
showMethods("toTable")
selectMethod("toTable", "Bimap")

```

---

type

*Accessing the type of an object*


---

**Description**

Get or set the *type* of an object.

Note that `type` and `type<-` are defined as *S4 generic functions* and what *type* means exactly (and what `type()` returns) depends on the objects for which the `type` and/or `type<-` methods are defined.

**Usage**

```

type(x)
type(x) <- value

## Methods defined in the BiocGenerics package:

## S4 method for signature 'vector'
type(x)
## S4 method for signature 'array'
type(x)
## S4 method for signature 'factor'
type(x) # returns "character"
## S4 method for signature 'data.frame'
type(x)

## S4 replacement method for signature 'vector'
type(x) <- value
## S4 replacement method for signature 'array'
type(x) <- value

```

**Arguments**

x	Any object for which the <code>type()</code> getter or setter is defined. Note that objects will either: not support the getter or setter at all, or support only the getter, or support both the getter and setter.
value	The type to set on x (assuming x supports the <code>type()</code> setter). value is typically (but not necessarily) expected to be a single string (i.e. a character vector of length 1).

## Details

On an ordinary vector, matrix, or array  $x$ , `type(x)` returns `typeof(x)`.

On a data frame  $x$  where all the columns are ordinary vectors or factors, `type(x)` is *semantically equivalent* to `typeof(as.matrix(x))`. However, the actual implementation is careful to avoid turning the full data frame  $x$  into a matrix, as this would tend to be very inefficient in general.

Note that for a matrix-like or array-like object, `type(x)` returns the type of the *elements* in the object. See `?S4Arrays::type` for more information.

## Value

`type(x)` is expected to return the type of  $x$  as a single string i.e. as a character vector of length 1.

## See Also

- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [type,ANY-method](#) in the **S4Arrays** package for the default type method.
- [type,DataFrame-method](#) in the **S4Arrays** package, and [type,PairwiseAlignments-method](#) in the **pwalign** package, for examples of specific type methods (defined for [DataFrame](#) and [PairwiseAlignments](#) objects, respectively).
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```

type
showMethods("type")

`type<-`
showMethods("type<-")

## The BiocGenerics package defines methods for ordinary vectors, arrays,
## and data frames:
m <- matrix(11:22, nrow=3)
type(m)          # equivalent to 'typeof(m)' or 'storage.mode(m)'
type(m) <- "raw" # equivalent to 'storage.mode(m) <- "raw"'
m
type(m)

selectMethod("type", "array")

selectMethod("type<-", "array")

df <- data.frame(a=44:49, b=letters[1:6], c=c(TRUE, FALSE))
stopifnot(identical(type(df), typeof(as.matrix(df))))

## Examples of methods defined in other packages:

library(S4Arrays)
showMethods("type")

```

```
selectMethod("type", "ANY") # the default "type" method

library(pwalign)
showMethods("type")
## The type() method for PairwiseAlignments objects:
selectMethod("type", "PairwiseAlignments")
```

---

unique

*Extract unique elements*


---

## Description

unique returns an object of the same class as `x` (typically a vector-like, data-frame-like, or array-like object) but with duplicate elements/rows removed.

NOTE: This man page is for the unique *S4 generic function* defined in the **BiocGenerics** package. See `?base::unique` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-like or data-frame-like) not supported by the default method.

## Usage

```
unique(x, incomparables=FALSE, ...)
```

## Arguments

`x` A vector-like, data-frame-like, or array-like object.  
`incomparables, ...`  
 See `?base::unique` for a description of these arguments.

## Value

See `?base::unique` for the value returned by the default method.

Specific methods defined in Bioconductor packages will typically return an object of the same class as the input object.

unique should always behave consistently with `BiocGenerics::duplicated`.

## See Also

- `base::unique` for the default unique method.
- `BiocGenerics::duplicated` for determining duplicate elements.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `unique,Rle-method` in the **S4Vectors** package for an example of a specific unique method (defined for **Rle** objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
unique
showMethods("unique")
selectMethod("unique", "ANY") # the default method
```

---

unlist

*Flatten list-like objects*


---

**Description**

Given a list-like object `x`, `unlist` produces a vector-like object obtained by concatenating (conceptually thru `c`) all the top-level elements in `x` (each of them being expected to be a vector-like object, typically).

NOTE: This man page is for the `unlist` *S4 generic function* defined in the **BiocGenerics** package. See `?base::unlist` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects not supported by the default method.

**Usage**

```
unlist(x, recursive=TRUE, use.names=TRUE)
```

**Arguments**

`x`                    A list-like object.  
`recursive, use.names`  
 See `?base::unlist` for a description of these arguments.

**Value**

See `?base::unlist` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

**See Also**

- `base::unlist` for the default `unlist` method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `unlist,List-method` in the **S4Vectors** package for an example of a specific `unlist` method (defined for `List` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
unlist # note the dispatch on the 'x' arg only
showMethods("unlist")
selectMethod("unlist", "ANY") # the default method
```

---

`unsplit`*Unsplit a list-like object*

---

### Description

Given a list-like object `value` and grouping `f`, `unsplit` produces a vector-like object `x` by conceptually reversing the split operation `value <- split(x, f)`.

NOTE: This man page is for the `unsplit` S4 generic function defined in the **BiocGenerics** package. See `?base::unsplit` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects not supported by the default method.

### Usage

```
unsplit(value, f, drop=FALSE)
```

### Arguments

<code>value</code>	A list-like object.
<code>f</code>	A factor or other grouping object that corresponds to the <code>f</code> symbol in <code>value &lt;- split(x, f)</code> .
<code>drop</code>	See <code>?base::unsplit</code> for a description of this argument.

### Value

See `?base::unsplit` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

### See Also

- `base::unsplit` for the default `unsplit` method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `unsplit,List-method` in the **IRanges** package for an example of a specific `unsplit` method (defined for `List` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

### Examples

```
unsplit # note the dispatch on the 'value' and 'f' args only
showMethods("unsplit")
selectMethod("unsplit", "ANY") # the default method
```

---

updateObject	<i>Update an object to its current class definition</i>
--------------	---

---

### Description

updateObject is a generic function that returns an instance of object updated to its current class definition.

### Usage

```
updateObject(object, ..., verbose=FALSE)

## Related utilities:
updateObjectFromSlots(object, objclass=class(object)[[1L]], ...,
                      verbose=FALSE)
getObjectSlots(object)
```

### Arguments

object	Object to be updated for updateObject and updateObjectFromSlots. Object for slot information to be extracted from for getObjectSlots.
...	Additional arguments, for use in specific updateObject methods.
verbose	TRUE or FALSE, indicating whether information about the update should be reported. Use <a href="#">message</a> to report this information.
objclass	Optional character string naming the class of the object to be created.

### Details

Updating objects is primarily useful when an object has been serialized (e.g., stored to disk) for some time (e.g., months), and the class definition has in the mean time changed. Because of the changed class definition, the serialized instance is no longer valid.

updateObject requires that the class of the returned object be the same as the class of the argument object, and that the object is valid (see [validObject](#)). By default, updateObject has the following behaviors:

updateObject(ANY, ..., verbose=FALSE) By default, updateObject uses heuristic methods to determine whether the object should be the 'new' S4 type (introduced in R 2.4.0), but is not. If the heuristics indicate an update is required, the updateObjectFromSlots function tries to update the object. The default method returns the original S4 object or the successfully updated object, or issues an error if an update is required but not possible. The optional named argument verbose causes a message to be printed describing the action. Arguments ... are passed to updateObjectFromSlots.

updateObject(list, ..., verbose=FALSE) Visit each element in list, applying updateObject(list[[elt]], ..., verbose=verbose).

updateObject(environment, ..., verbose=FALSE) Visit each element in environment, applying updateObject(environment[[elt]], ..., verbose=verbose)

`updateObject(formula, ..., verbose=FALSE)` Do nothing; the environment of the formula may be too general (e.g., `R_GlobalEnv`) to attempt an update.

`updateObject(envRefClass, ..., verbose=FALSE)` Attempt to update objects from fields using a strategy like `updateObjectFromSlots Method 1`.

`updateObjectFromSlots(object, objclass=class(object), ..., verbose=FALSE)` is a utility function that identifies the intersection of slots defined in the object instance and `objclass` definition. Under Method 1, the corresponding elements in `object` are then updated (with `updateObject(elt, ..., verbose=verbose)`) and used as arguments to a call to `new(class, ...)`, with `...` replaced by slots from the original object. If this fails, then Method 2 tries `new(class)` and assigns slots of object to the newly created instance.

`getObjectSlots(object)` extracts the slot names and contents from `object`. This is useful when `object` was created by a class definition that is no longer current, and hence the contents of `object` cannot be determined by accessing known slots.

## Value

`updateObject` returns a valid instance of `object`.

`updateObjectFromSlots` returns an instance of class `objclass`.

`getObjectSlots` returns a list of named elements, with each element corresponding to a slot in `object`.

## See Also

- [updateObjectTo](#) in the **Biobase** package for updating an object to the class definition of a template (might be useful for updating a virtual superclass).
- [validObject](#) for testing the validity of an object.
- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
updateObject
showMethods("updateObject")
selectMethod("updateObject", "ANY") # the default method

library(Biobase)
## update object, same class
data(sample.ExpressionSet)
obj <- updateObject(sample.ExpressionSet)

setClass("UpdtA", representation(x="numeric"), contains="data.frame")
setMethod("updateObject", "UpdtA",
  function(object, ..., verbose=FALSE)
  {
    if (verbose)
      message("updateObject object = 'A'")
```

```

        object <- callNextMethod()
        object@x <- -object@x
        object
    }
)

a <- new("UpdtA", x=1:10)
## See steps involved
updateObject(a)

removeMethod("updateObject", "UpdtA")
removeClass("UpdtA")

```

var

---

### Variance and Standard Deviation

---

**Description**

`var` and `sd` compute the variance and standard deviation of a vector `x`.

NOTE: This man page is for the `var` and `sd`, *S4 generic functions* defined in the **BiocGenerics** package. See `?stats::var` and `?stats::sd` for the default methods (defined in the **stats** package). Bioconductor packages can define specific methods for objects (typically array-like) not supported by the default method.

**Usage**

```

var(x, y = NULL, na.rm = FALSE, use)
sd(x, na.rm = FALSE)

```

**Arguments**

<code>x</code>	a vector-like object
<code>y</code>	a vector-like object, or <code>NULL</code>
<code>na.rm, use</code>	see <a href="#">var</a>

**Value**

See `?stats::var` and `?stats::sd` for the value returned by the default methods.

Specific methods defined in Bioconductor packages will typically return an object of the same class as the input object.

**See Also**

- `stats::var` and `stats::sd` for the default methods.
- [showMethods](#) for displaying a summary of the methods defined for a given generic function.
- [selectMethod](#) for getting the definition of a specific method.
- [BiocGenerics](#) for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
var
showMethods("var")
selectMethod("var", "ANY") # the default method
```

---

weights

*Extract model weights*

---

## Description

weights is a generic function which extracts fitting weights from objects returned by modeling functions.

NOTE: This man page is for the weights *S4 generic function* defined in the **BiocGenerics** package. See `?stats::weights` for the default method (defined in the **stats** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```
weights(object, ...)
```

## Arguments

object, ... See `?stats::weights`.

## Value

Weights extracted from the object object.

See `?stats::weights` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## See Also

- `stats::weights` for the default weights method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `weights,PLMset-method` in the **affyPLM** package for an example of a specific weights method (defined for `PLMset` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
weights
showMethods("weights")
selectMethod("weights", "ANY") # the default method
```

---

which

*Which values in an object are considered TRUE?*

---

## Description

Give the indices of the values in a vector-, array-, or list-like object that are considered TRUE, allowing for array indices in the case of an array-like object.

NOTE: This man page is for the `which` *S4 generic function* defined in the **BiocGenerics** package. See `?base::which` for the default method (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-, array-, or list-like) not supported by the default methods.

## Usage

```
which(x, arr.ind=FALSE, useNames=TRUE)
```

## Arguments

`x` An object, typically with a vector-, array-, or list-like semantic.  
`arr.ind, useNames`  
See `?base::which` for a description of these arguments.

## Value

See `?base::which` for the value returned by the default method.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default method.

## See Also

- `base::which` for the default `which` method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `which,DelayedArray-method` in the **DelayedArray** package for an example of a specific `which` method (defined for **DelayedArray** objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```
which
showMethods("which")
selectMethod("which", "ANY") # the default method

library(DelayedArray)
showMethods("which")
## The which() method for DelayedArray objects:
selectMethod("which", "DelayedArray")
```

---

`which.min`*What's the index of the first min or max value in an object?*

---

## Description

Determines the location (i.e. index) of the (first) minimum or maximum value in an object.

NOTE: This man page is for the `which.min` and `which.max` *S4 generic functions* defined in the **BiocGenerics** package. See `?base::which.min` for the default methods (defined in the **base** package). Bioconductor packages can define specific methods for objects (typically vector-, array-, or list-like) not supported by the default methods.

## Usage

```
which.min(x, ...)  
which.max(x, ...)
```

## Arguments

<code>x</code>	An object, typically with a vector-, array-, or list-like semantic.
<code>...</code>	Additional arguments, for use in specific methods.

## Value

See `?base::which.min` for the value returned by the default methods.

Specific methods defined in Bioconductor packages should behave as consistently as possible with the default methods.

## Note

The default methods (defined in the **base** package) only take a single argument. We've added the `...` argument to the generic functions defined in the **BiocGenerics** package so they can be called with an arbitrary number of effective arguments. This typically allows methods to add extra arguments for controlling/altering the behavior of the operation. Like for example the global argument supported by the `which.max` method for **NumericList** objects (defined in the **IRanges** package).

## See Also

- `base::which.min` for the default `which.min` and `which.max` methods.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `which.max,NumericList-method` in the **IRanges** package for an example of a specific `which.max` method (defined for **NumericList** objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

## Examples

```

which.min
showMethods("which.min")
selectMethod("which.min", "ANY") # the default method

which.max
showMethods("which.max")
selectMethod("which.max", "ANY") # the default method

library(IRanges)
showMethods("which.max")
## The which.max() method for NumericList objects:
selectMethod("which.max", "NumericList")

```

---

xtabs

*Cross tabulation*


---

## Description

xtabs creates a contingency table (optionally a sparse matrix) from cross-classifying factors, usually contained in a data-frame-like object, using a formula interface.

NOTE: This man page is for the xtabs *S4 generic function* defined in the **BiocGenerics** package. See `?stats::xtabs` for the default method (defined in the **stats** package). Bioconductor packages can define specific methods for objects not supported by the default method.

## Usage

```

xtabs(formula=~., data=parent.frame(), subset, sparse=FALSE,
      na.action, na.rm=FALSE, addNA=FALSE, exclude=if(!addNA)c(NA, NaN),
      drop.unused.levels=FALSE)

```

## Arguments

formula, subset, sparse, na.action, na.rm, addNA, exclude,  
drop.unused.levels

See `?stats::xtabs` for a description of these arguments.

data            A data-frame-like object.

## Value

See `?stats::xtabs` for the value returned by the default method.

Specific methods defined in Bioconductor packages should also return the type of object returned by the default method.

**See Also**

- `stats::xtabs` for the default xtabs method.
- `showMethods` for displaying a summary of the methods defined for a given generic function.
- `selectMethod` for getting the definition of a specific method.
- `xtabs,DataFrame-method` in the **S4Vectors** package for an example of a specific xtabs method (defined for `DataFrame` objects).
- **BiocGenerics** for a summary of all the generics defined in the **BiocGenerics** package.

**Examples**

```
xtabs # note the dispatch on the 'data' arg only
showMethods("xtabs")
selectMethod("xtabs", "ANY") # the default method

library(S4Vectors)
showMethods("xtabs")
## The xtabs() method for DataFrame objects:
selectMethod("xtabs", "DataFrame")
```

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