

# Package ‘ff’

May 15, 2018

**Version** 2.2-14

**Date** 2018-04-15

**Title** Memory-Efficient Storage of Large Data on Disk and Fast Access Functions

**Author** Daniel Adler <dadler@uni-goettingen.de>,  
Christian Gläser <christian\_glaeser@gmx.de>,  
Oleg Nenadic <onenadi@uni-goettingen.de>,  
Jens Oehlschlägel <Jens.Oehlschlaegel@truecluster.com>,  
Walter Zucchini <wzucchi@uni-goettingen.de>

**Maintainer** Jens Oehlschlägel <Jens.Oehlschlaegel@truecluster.com>

**Depends** R (>= 2.10.1), bit (>= 1.1-13), utils

**Suggests** biglm

**Description** The ff package provides data structures that are stored on disk but behave (almost) as if they were in RAM by transparently mapping only a section (pagesize) in main memory - the effective virtual memory consumption per ff object. ff supports R's standard atomic data types 'double', 'logical', 'raw' and 'integer' and non-standard atomic types boolean (1 bit), quad (2 bit unsigned), nibble (4 bit unsigned), byte (1 byte signed with NAs), ubyte (1 byte unsigned), short (2 byte signed with NAs), ushort (2 byte unsigned), single (4 byte float with NAs). For example 'quad' allows efficient storage of genomic data as an 'A','T','G','C' factor. The unsigned types support 'circular' arithmetic. There is also support for close-to-atomic types 'factor', 'ordered', 'POSIXct', 'Date' and custom close-to-atomic types.

ff not only has native C-support for vectors, matrices and arrays with flexible dimorder (major column-order, major row-order and generalizations for arrays). There is also a ffd class not unlike data.frames and import/export filters for csv files.

ff objects store raw data in binary flat files in native encoding, and complement this with metadata stored in R as physical and virtual attributes. ff objects have well-defined hybrid copying semantics, which gives rise to certain performance improvements through virtualization. ff objects can be stored and reopened across R

sessions. ff files can be shared by multiple ff R objects (using different data en/de-coding schemes) in the same process or from multiple R processes to exploit parallelism. A wide choice of finalizer options allows to work with 'permanent' files as well as creating/removing 'temporary' ff files completely transparent to the user. On certain OS/Filesystem combinations, creating the ff files works without notable delay thanks to using sparse file allocation. Several access optimization techniques such as Hybrid Index Preprocessing and Virtualization are implemented to achieve good performance even with large datasets, for example virtual matrix transpose without touching a single byte on disk. Further, to reduce disk I/O, 'logicals' and non-standard data types get stored native and compact on binary flat files i.e. logicals take up exactly 2 bits to represent TRUE, FALSE and NA.

Beyond basic access functions, the ff package also provides compatibility functions that facilitate writing code for ff and ram objects and support for batch processing on ff objects (e.g. as.ram, as.ff, ffapply). ff interfaces closely with functionality from package 'bit': chunked looping, fast bit operations and coercions between different objects that can store subscript information ('bit', 'bitwhich', ff 'boolean', ri range index, hi hybrid index). This allows to work interactively with selections of large datasets and quickly modify selection criteria.

Further high-performance enhancements can be made available upon request.

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**LazyLoad** yes

**ByteCompile** yes

**Encoding** latin1

**URL** <http://ff.r-forge.r-project.org/>

**NeedsCompilation** yes

**Repository** CRAN

**Date/Publication** 2018-05-15 21:19:05 UTC

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---

**add** *Incrementing an ff or ram object*

---

**Description**

Yet another assignment interface in order to allow to formulate `x[index, ..., add=TRUE]<-value` in a way which works transparently, not only for ff, but also for ram objects: `add(x, value, index, ...)`.

**Usage**

```
add(x, ...)
## S3 method for class 'ff'
add(x, value, ...)
## Default S3 method:
add(x, value, ...)
```

**Arguments**

<code>x</code>	an ff or ram object
<code>value</code>	the amount to increment, possibly recycled
<code>...</code>	further arguments – especially index information – passed to <code>[&lt;-</code> or <code>[&lt;- .ff</code>

**Value**

`invisible()`

**Note**

Note that `add.default` changes the object in its parent frame and thus violates R's usual functional programming logic. Duplicated index positions should be avoided, because ff and ram objects behave differently:

```
add.ff(x, 1, c(3,3))
# will increment x at position 3 TWICE by 1, while
add.default(x, 1, c(3,3))
# will increment x at position 3 just ONCE by 1
```

**Author(s)**

Jens Oehlschlägel

**See Also**

[swap](#), [\[.ff](#), [LimWarn](#)

**Examples**

```

message("incrementing parts of a vector")
x <- ff(0, length=12)
y <- rep(0, 12)
add(x, 1, 1:6)
add(y, 1, 1:6)
x
y

message("incrementing parts of a matrix")
x <- ff(0, dim=3:4)
y <- array(0, dim=3:4)
add(x, 1, 1:2, 1:2)
add(y, 1, 1:2, 1:2)
x
y

message("BEWARE that ff and ram methods differ in treatment of duplicated index positions")
add(x, 1, c(3,3))
add(y, 1, c(3,3))
x
y

rm(x); gc()

```

---

array2vector

*Array: make vector from array*


---

**Description**

Makes a vector from an array respecting 'dim' and 'dimorder'

**Usage**

```
array2vector(x, dim = NULL, dimorder = NULL)
```

**Arguments**

x	an <a href="#">array</a>
dim	<a href="#">dim</a>
dimorder	<a href="#">dimorder</a>

**Details**

This is the inverse function of [vector2array](#). It extracts the vector from the array by first moving through the fastest rotating dimension `dim[dimorder[1]]`, then `dim[dimorder[2]]`, and so forth

**Value**

a vector

**Author(s)**

Jens Oehlschlägel

**See Also**

[vector2array](#), [arrayIndex2vectorIndex](#)

**Examples**

```
array2vector(matrix(1:12, 3, 4))
array2vector(matrix(1:12, 3, 4, byrow=TRUE), dimorder=2:1)
```

---

arrayIndex2vectorIndex

*Array: make vector positions from array index*

---

**Description**

Make vector positions from a (non-symmetric) array index respecting 'dim' and 'dimorder'

**Usage**

```
arrayIndex2vectorIndex(x, dim = NULL, dimorder = NULL, vw = NULL)
```

**Arguments**

x	an n by m matrix with n m-dimensional array indices
dim	NULL or <a href="#">dim</a>
dimorder	NULL or <a href="#">dimorder</a>
vw	NULL or integer vector[3] or integer matrix[3,m], see details

**Details**

The fastest rotating dimension is dim[dimorder[1]], then dim[dimorder[2]], and so forth. The parameters 'x' and 'dim' may refer to a subarray of a larger array, in this case, the array indices 'x' are interpreted as 'vw[1,] + x' within the larger array 'as.integer(colSums(vw))'.

**Value**

a vector of indices in 1:prod(dim) (or 1:prod(colSums(vw)))

**Author(s)**

Jens Oehlschlägel

**See Also**

[array2vector](#), [vectorIndex2arrayIndex](#)

**Examples**

```
x <- matrix(1:12, 3, 4)
x
arrayIndex2vectorIndex(cbind(as.vector(row(x)), as.vector(col(x)))
, dim=dim(x))
arrayIndex2vectorIndex(cbind(as.vector(row(x)), as.vector(col(x)))
, dim=dim(x), dimorder=2:1)
matrix(1:30, 5, 6)
arrayIndex2vectorIndex(cbind(as.vector(row(x)), as.vector(col(x)))
, vw=rbind(c(0,1), c(3,4), c(2,1)))
arrayIndex2vectorIndex(cbind(as.vector(row(x)), as.vector(col(x)))
, vw=rbind(c(0,1), c(3,4), c(2,1)), dimorder=2:1)
```

---

as.ff

*Coercing ram to ff and ff to ram objects*


---

**Description**

Coercing ram to ff and ff to ram objects while optionally modifying object features.

**Usage**

```
as.ff(x, ...)
as.ram(x, ...)
## Default S3 method:
as.ff(x, filename = NULL, overwrite = FALSE, ...)
## S3 method for class 'ff'
as.ff(x, filename = NULL, overwrite = FALSE, ...)
## Default S3 method:
as.ram(x, ...)
## S3 method for class 'ff'
as.ram(x, ...)
```

**Arguments**

x	any object to be coerced
filename	path and filename
overwrite	TRUE to overwrite the old filename
...	...



## Details

If `as.ff.ff` is called on an 'ff' object or `as.ram.default` is called on a non-ff object AND no changes are required, the input object 'x' is returned unchanged. Otherwise the workhorse `clone.ff` is called. If no change of features are requested, the filename attached to the object remains unchanged, otherwise a new filename is requested (or can be set by the user).

## Value

A ram or ff object.

## Note

If you use `ram <- as.ram(ff)` for caching, please note that you must `close.ff` before you can write back `as.ff(ram, overwrite=TRUE)` (see examples).

## Author(s)

Jens Oehlschlägel

## See Also

[as.ff.bit](#), [ff](#), [clone](#), [as.vmode](#), [vmode](#), [as.hi](#)

## Examples

```
message("create ff")
myintff <- ff(1:12)
message("coerce (=clone) integer ff to double ff")
mydoubleff <- as.ff(myintff, vmode="double")
message("cache (=clone) integer ff to integer ram AND close original ff")
myintram <- as.ram(myintff) # filename is retained
close(myintff)
message("modify ram cache and write back (=clone) to ff")
myintram[1] <- -1L
myintff <- as.ff(myintram, overwrite=TRUE)
message("coerce (=clone) integer ram to double ram")
mydoubleram <- as.ram(myintram, vmode="double")
message("coerce (inplace) integer ram to double ram")
myintram <- as.ram(myintram, vmode="double")
message("more classic: coerce (inplace) double ram to integer ram")
vmode(myintram) <- "integer"
rm(myintff, myintram, mydoubleff, mydoubleram); gc()
```

---

`as.ff.bit`*Conversion between bit and ff boolean*

---

### Description

Function `as.ff.bit` converts a `bit` vector to a boolean `ff` vector. Function `as.bit.ff` converts a boolean `ff` vector to a `ff` vector.

### Usage

```
## S3 method for class 'bit'  
as.ff(x, filename = NULL, overwrite = FALSE, ...)  
## S3 method for class 'ff'  
as.bit(x, ...)
```

### Arguments

<code>x</code>	the source of conversion
<code>filename</code>	optionally a desired filename
<code>overwrite</code>	logical indicating whether we allow overwriting the target file
<code>...</code>	further arguments passed to <code>ff</code> in case <code>as.ff.bit</code> , ignored in case of <code>as.bit.ff</code>

### Details

The data are copied bot bit-wise but integerwise, therefore these conversions are very fast. `as.bit.ff` will attach the `ff` filename to the bit vector, and `as.ff.bit` will - if attached - use THIS filename and SILENTLY overwrite this file.

### Value

A vector of the converted type

### Note

NAs are mapped to TRUE in 'bit' and to FALSE in 'ff' booleans. Might be aligned in a future release. Don't use bit if you have NAs - or map NAs explicetely.

### Author(s)

Jens Oehlschlägel

### See Also

[bit](#), [ff](#), [as.ff](#), [as.hi.bit](#)

**Examples**

```

l <- as.boolean(sample(c(FALSE,TRUE), 1000, TRUE))

b <- as.bit(l)
stopifnot(identical(l,b[]))
b
f <- as.ff(b)
stopifnot(identical(l,f[]))
f
b2 <- as.bit(f)
stopifnot(identical(l,b2[]))
b2
f2 <- as.ff(b2)
stopifnot(identical(filename(f),filename(f2)))
stopifnot(identical(l,f2[]))
f
rm(f,f2); gc()

```

as.ffdf

*Coercing to ffdf and data.frame***Description**

Functions for coercing to ffdf and data.frame

**Usage**

```

as.ffdf(x, ...)
## S3 method for class 'ff_vector'
as.ffdf(x, ...)
## S3 method for class 'ff_matrix'
as.ffdf(x, ...)
## S3 method for class 'data.frame'
as.ffdf(x, vmode=NULL, col_args = list(), ...)
## S3 method for class 'ffdf'
as.data.frame(x, ...)

```

**Arguments**

x	the object to be coerced
vmode	optional specification of the <code>vmodes</code> of columns of the <code>data.frame</code> . Either a character vector of <code>vmodes</code> (named with column names of the <code>data.frame</code> or recycled if not named) or a list named with <code>vmodes</code> where each element identifies those columns of the <code>data.frame</code> that should get the <code>vmode</code> encoded in the name of the element
col_args	further arguments; passed to <code>ff</code>
...	further arguments; passed to <code>ffdf</code> for <code>.ff_vector</code> , <code>.ff_matrix</code> and <code>.data.frame</code> methods, ignored for <code>.ffdf</code> identity method

**Value**

'as.ffdf' returns an object of class `ffdf`, 'as.data.frame' returns an object of class `data.frame`

**Author(s)**

Jens Oehlschlägel

**See Also**

[is.ffdf](#), [ffdf](#), [data.frame](#)

**Examples**

```
d <- data.frame(x=1:26, y=letters, z=Sys.time()+1:26)
ffd <- as.ffdf(d)
stopifnot(identical(d, as.data.frame(ffd)))
rm(ffd); gc()
```

---

as.hi

*Hybrid Index, coercion to*

---

**Description**

The generic `as.hi` and its methods are the main (internal) means for preprocessing index information into the hybrid index class `hi`. Usually `as.hi` is called transparently from `[.ff`. However, you can explicitly do the index-preprocessing, store the Hybrid Index `hi`, and use the `hi` for subscripting.

**Usage**

```
as.hi(x, ...)
## S3 method for class 'hi'
as.hi(x, ...)
## S3 method for class 'ri'
as.hi(x, maxindex = length(x), ...)
## S3 method for class 'bit'
as.hi(x, range = NULL, maxindex = length(x), vw = NULL
, dim = NULL, dimorder = NULL, pack = TRUE, ...)
## S3 method for class 'bitwhich'
as.hi(x, maxindex = length(x), pack = FALSE, ...)
## S3 method for class 'call'
as.hi(x, maxindex = NA, dim = NULL, dimorder = NULL, vw = NULL
, vw.convert = TRUE, pack = TRUE, envir = parent.frame(), ...)
## S3 method for class 'name'
as.hi(x, envir = parent.frame(), ...)

## S3 method for class 'integer'
```

```

as.hi(x, maxindex = NA, dim = NULL, dimorder = NULL
, symmetric = FALSE, fixdiag = NULL, vw = NULL, vw.convert = TRUE
, dimorder.convert = TRUE, pack = TRUE, NAs = NULL, ...)
## S3 method for class 'which'
as.hi(x, ...)
## S3 method for class 'double'
as.hi(x, ...)
## S3 method for class 'logical'
as.hi(x, maxindex = NA, dim = NULL, vw = NULL, pack = TRUE, ...)
## S3 method for class 'character'
as.hi(x, names, vw = NULL, vw.convert = TRUE, ...)
## S3 method for class 'matrix'
as.hi(x, dim, dimorder = NULL, symmetric = FALSE, fixdiag = NULL
, vw = NULL, pack = TRUE, ...)

```

### Arguments

x	an appropriate object of the class for which we dispatched
envir	the environment in which to evaluate components of the index expression
maxindex	maximum positive indexposition maxindex, is needed with negative indices, if vw or dim is given, maxindex is calculated automatically
names	the <code>names</code> of the indexed vector for character indexing
dim	the <code>dim</code> of the indexed matrix to be stored within the <code>hi</code> object
dimorder	the <code>dimorder</code> of the indexed matrix to be stored within the <code>hi</code> object, may convert interpretation of x
symmetric	the <code>symmetric</code> of the indexed matrix to be stored within the <code>hi</code> object
fixdiag	the <code>fixdiag</code> of the indexed matrix to be stored within the <code>hi</code> object
vw	the virtual window <code>vw</code> of the indexed vector or matrix to be stored within the <code>hi</code> object, see details
vw.convert	FALSE to prevent doubly virtual window conversion, this is needed for some internal calls that have done the virtual window conversion already, see details
dimorder.convert	FALSE to prevent doubly dimorder conversion, this is needed for some internal calls that have done the dimorder conversion already, see details
NAs	a vector of NA positions to be stored <code>rlepacked</code> , not fully supported yet
pack	FALSE to prevent <code>rlepacking</code> , note that this is a hint rather than a guarantee, <code>as.hi.bit</code> might ignore this
range	NULL or a vector with two elements indicating first and last position to be converted from 'bit' to 'hi'
...	further argument passed from generic to method or from wrapper method to <code>as.hi.integer</code>

## Details

The generic dispatches appropriately, `as.hi.hi` returns an `hi` object unchanged, `as.hi.call` tries to `hiparse` instead of evaluate its input in order to save RAM. If parsing is successful `as.hi.call` will ignore its argument `pack` and always `pack` unless the subscript is too small to do so. If parsing fails it evaluates the index expression and dispatches again to one of the other methods. `as.hi.name` and `as.hi.(` are wrappers to `as.hi.call`. `as.hi.integer` is the workhorse for coercing evaluated expressions and `as.hi.which` is a wrapper removing the `which` class attribute. `as.hi.double`, `as.hi.logical` and `as.hi.character` are also wrappers to `as.hi.integer`, but note that `as.hi.logical` is not memory efficient because it expands *all* positions and then applies logical subscripting.

`as.hi.matrix` calls `arrayIndex2vectorIndex` and then `as.hi.integer` to interpret and preprocess matrix indices.

If the `dim` and `dimorder` parameter indicate a non-standard dimorder (`dimorderStandard`), the index information in `x` is converted from a standard dimorder interpretation to the requested `dimorder`. If the `vw` parameter is used, the index information in `x` is interpreted relative to the virtual window but stored relative to the absolute origin. Back-coercion via `as.integer.hi` and friends will again return the index information relative to the virtual window, thus retaining symmetry and transparency of the virtual window to the user.

You can use `length` to query the index length (possibly length of negative subscripts), `poslength` to query the number of selected elements (even with negative subscripts), and `maxindex` to query the largest possible index position (within virtual window, if present)

Duplicated negative indices are removed and will not be recovered by `as.integer.hi`.

## Value

an object of class `hi`

## Note

Avoid changing the Hybrid Index representation, this might crash the `[.ff` subscripting.

## Author(s)

Jens Oehlschlägel

## See Also

`hi` for the Hybrid Index class, `hiparse` for parsing details, `as.integer.hi` for back-coercion, `[.ff` for `ff` subscripting

## Examples

```
message("integer indexing with and without rel-packing")
as.hi(1:12)
as.hi(1:12, pack=FALSE)
message("if index is double, the wrapper method just converts to integer")
as.hi(as.double(1:12))
message("if index is character, the wrapper method just converts to integer")
as.hi(c("a","b","c"), names=letters)
message("negative index must use maxindex (or vw)")
```

```

as.hi(-(1:3), maxindex=12)
message("logical index can use maxindex")
as.hi(c(FALSE, FALSE, TRUE, TRUE))
as.hi(c(FALSE, FALSE, TRUE, TRUE), maxindex=12)

message("matrix index")
x <- matrix(1:12, 6)
as.hi(rbind(c(1,1), c(1,2), c(2,1)), dim=dim(x))

message("first ten positions within virtual window")
i <- as.hi(1:10, vw=c(10, 80, 10))
i
message("back-coerce relativ to virtual window")
as.integer(i)
message("back-coerce relativ to absolute origin")
as.integer(i, vw.convert=FALSE)

message("parsed index expressions save index RAM")
as.hi(quote(1:100000000))
## Not run:
message("compare to RAM requirement when the index experssion is evaluated")
as.hi(1:100000000)

## End(Not run)

message("example of parsable index expression")
a <- seq(100, 200, 20)
as.hi(substitute(c(1:5, 4:9, a)))
hi(c(1,4, 100),c(5,9, 200), by=c(1,1,20))

message("two examples of index expression temporarily expanded to full length due to
non-supported use of brackets '(' and mathematical operators '+' accepting token")
message("example1: accepted token but aborted parsing because length>16")
as.hi(quote(1+(1:16)))
message("example1: rejected token and aborted parsing because length>16")
as.hi(quote(1+(1:17)))

```

---

as.integer.hi

*Hybrid Index, coercing from*


---

## Description

Functions that (back-)convert an `hi` object to the respective subscripting information.

## Usage

```

## S3 method for class 'hi'
as.which(x, ...)
## S3 method for class 'hi'
as.bitwhich(x, ...)

```

```
## S3 method for class 'hi'
as.bit(x, ...)
## S3 method for class 'hi'
as.integer(x, vw.convert = TRUE, ...)
## S3 method for class 'hi'
as.logical(x, maxindex = NULL, ...)
## S3 method for class 'hi'
as.character(x, names, vw.convert = TRUE, ...)
## S3 method for class 'hi'
as.matrix(x, dim = x$dim, dimorder = x$dimorder
, vw = x$vw, symmetric = x$symmetric, fixdiag = x$fixdiag, ...)
```

### Arguments

x	an object of class <a href="#">hi</a>
maxindex	the <a href="#">length</a> of the subscripted object (needed for logical output)
names	the <a href="#">names</a> vector of the subscripted object
dim	the <a href="#">dim</a> of the subscripted object
dimorder	the <a href="#">dimorder</a> of the subscripted object
vw	the virtual window <a href="#">vw</a> of the subscripted object
vw.convert	vw.convert
symmetric	TRUE if the subscripted matrix is <a href="#">symmetric</a>
fixdiag	TRUE if the subscripted matrix has <a href="#">fixdiag</a>
...	further arguments passed

### Value

as.integer.hi returns an integer vector, see [as.hi.integer](#). as.logical.hi returns an logical vector, see [as.hi.logical](#). as.character.hi returns a character vector, see [as.hi.character](#). as.matrix.hi returns a matrix index, see [as.hi.matrix](#).

### Author(s)

Jens Oehlschlägel

### See Also

[hi](#), [as.hi](#)

### Examples

```
x <- 1:6
names(x) <- letters[1:6]
as.integer(as.hi(c(1:3)))
as.logical(as.hi(c(TRUE,TRUE,TRUE,FALSE,FALSE,FALSE)))
as.character(as.hi(letters[1:3], names=names(x)), names=names(x))
x <- matrix(1:12, 6)
as.matrix(as.hi(rbind(c(1,1), c(1,2), c(2,1)), dim=dim(x)), dim=dim(x))
```



**Description**

as.vmode is a generic that converts some R ram object to the desired [vmode](#).

**Usage**

```
as.vmode(x, ...)
as.boolean(x, ...)
as.quad(x, ...)
as.nibble(x, ...)
as.byte(x, ...)
as.ubyte(x, ...)
as.short(x, ...)
as.ushort(x, ...)
## Default S3 method:
as.vmode(x, vmode, ...)
## S3 method for class 'ff'
as.vmode(x, ...)
## Default S3 method:
as.boolean(x, ...)
## Default S3 method:
as.quad(x, ...)
## Default S3 method:
as.nibble(x, ...)
## Default S3 method:
as.byte(x, ...)
## Default S3 method:
as.ubyte(x, ...)
## Default S3 method:
as.short(x, ...)
## Default S3 method:
as.ushort(x, ...)
```

**Arguments**

x	any object
vmode	virtual mode
...	The ... don't have a function yet, they are only defined to keep the generic flexible.

**Details**

Function as.vmode actually coerces to one of the usual [storage.modes](#) (see [.rammode](#)) but flags them with an additional attribute 'vmode' if necessary. The coercion generics can also be called

directly:

as.boolean	1 bit logical without NA
as.logical	2 bit logical with NA
as.quad	2 bit unsigned integer without NA
as.nibble	4 bit unsigned integer without NA
as.byte	8 bit signed integer with NA
as.ubyte	8 bit unsigned integer without NA
as.short	16 bit signed integer with NA
as ushort	16 bit unsigned integer without NA
as.integer	32 bit signed integer with NA
as.single	32 bit float
as.double	64 bit float
as.complex	2x64 bit float
as.raw	8 bit unsigned char
as.character	character

### Value

a vector of the desired vmode containing the input data

### Author(s)

Jens Oehlschlägel

### See Also

[vmode](#), [vector.vmode](#)

### Examples

```
as.vmode(1:3,"double")
as.vmode(1:3,"byte")
as.double(1:3)
as.byte(1:3)
```

---

bigsample

*Sampling from large pools*

---

### Description

bigsample samples quicker from large pools than [sample](#) does.

**Usage**

```
bigsample(x, ...)
## Default S3 method:
bigsample(x, size, replace = FALSE, prob = NULL, negative = FALSE, ...)
## S3 method for class 'ff'
bigsample(x, size, replace = FALSE, prob = NULL, ...)
```

**Arguments**

x	the pool to sample from
size	the number of elements to sample
replace	TRUE to use sampling with replacement
prob	optional vector of sampling probabilities (recycled to pool length)
negative	negative
...	...

**Details**

For small pools [sample](#) is called.

**Value**

a vector of elements sampled from the pool (argument 'x')

**Note**

Note that `bigsample` and `sample` do not necessarily return the same sequence of elements when `set.seed` is set before.

**Author(s)**

Daniel Adler, Jens Oehlschlägel, Walter Zucchini

**See Also**

[sample](#), [ff](#)

**Examples**

```
message("Specify pool size")
bigsample(1e8, 10)
message("Sample ff elements (same as x[bigsample(length(ff(1:100 / 10)), 10)])")
bigsample(ff(1:100 / 10), 10)
## Not run:
message("Speed factor")
(system.time(for(i in 1:10)sample(1e8, 10))[3]/10)
/ (system.time(for(i in 1:1000)bigsample(1e8, 10))[3]/1000)

## End(Not run)
```

**Description**

These are used in aggregating the chunks resulting from batch processing. They are usually called via [do.call](#)

**Usage**

```
ccbind(...)
crbind(...)
cfun(..., FUN, FUNARGS = list())
cquantile(..., probs = seq(0, 1, 0.25), na.rm = FALSE, names = TRUE, type = 7)
csummary(..., na.rm = "ignored")
cmedian(..., na.rm = FALSE)
clength(..., na.rm = FALSE)
csum(..., na.rm = FALSE)
cmean(..., na.rm = FALSE)
```

**Arguments**

...	...
FUN	a aggregating function
FUNARGS	further arguments to the aggregating function
na.rm	TRUE to remove NAs
probs	see <a href="#">quantile</a>
names	see <a href="#">quantile</a>
type	see <a href="#">quantile</a>

**Details**

CFUN	FUN	comment
ccbind	<a href="#">cbind</a>	like cbind but respecting names
crbind	<a href="#">rbind</a>	like rbind but respecting names
cfun		crbind the input chunks and then apply 'FUN' to each column
cquantile	<a href="#">quantile</a>	crbind the input chunks and then apply 'quantile' to each column
csummary	<a href="#">summary</a>	crbind the input chunks and then apply 'summary' to each column
cmedian	<a href="#">median</a>	crbind the input chunks and then apply 'median' to each column
clength	<a href="#">length</a>	crbind the input chunks and then determine the number of values in each column
csum	<a href="#">sum</a>	crbind the input chunks and then determine the sum values in each column
cmean	<a href="#">mean</a>	crbind the input chunks and then determine the (unweighted) mean in each column

In order to use CFUNs on the result of `lapply` or `ffapply` use `do.call`.

### Value

depends on the CFUN used

### ff options

xx TODO: extend this for weighted means, weighted median etc.,  
see <http://tolstoy.newcastle.edu.au/R/help/02a/1073.html> and <http://tolstoy.newcastle.edu.au/R/help/02a/1060.html>  
or google "Re: [R] Weighted median"

### Note

Currently - for command line convenience - we map the elements of a single list argument to ..., but this may change in the future.

### Author(s)

Jens Oehlschlägel

### See Also

[ffapply](#), [do.call](#), [na.count](#)

### Examples

```
X <- lapply(split(rnorm(1000), 1:10), summary)
do.call("crbind", X)
do.call("csummary", X)
do.call("cmean", X)
do.call("cfun", c(X, list(FUN=mean, FUNARGS=list(na.rm=TRUE))))
rm(X)
```

---

chunk.bit

*Chunk bit vectors*

---

### Description

chunking method for ffd objects automatically considering RAM requirements from recordsize as calculated from `sum(.rambytes[vmode])`

### Usage

```
## S3 method for class 'bit'
chunk(x, RECORDBYTES = .rambytes["logical"], BATCHBYTES = getOption("ffbatchbytes"), ...)
```

**Arguments**

x	<a href="#">bit</a>
RECORDBYTES	optional integer scalar representing the bytes needed to process a single element of the bit vector
BATCHBYTES	integer scalar limiting the number of bytes to be processed in one chunk, default from <code>getOption("ffbatchbytes")</code> , see also <a href="#">.rambytes</a>
...	further arguments passed to <a href="#">chunk</a>

**Value**

A list with [ri](#) indexes each representing one chunk

**Author(s)**

Jens Oehlschlägel

**See Also**

[chunk](#), [bit](#)

**Examples**

```
n <- 1000
x <- bit(n)
ceiling(n / (300 %% sum(.rambytes["logical"])))
chunk(x, BATCHBYTES=300)
ceiling((n/2) / (100 %% sum(.rambytes["logical"])))
chunk(x, from=1, to = n/2, BATCHBYTES=100)
rm(x, n)
```

---

chunk.ffdf

*Chunk\_ff\_vector and ffdf*

---

**Description**

Chunking method for `ff_vector` and `ffdf` objects (row-wise) automatically considering RAM requirements from `recordsize` as calculated from `sum(.rambytes[vmode])`

**Usage**

```
## S3 method for class 'ff_vector'
chunk(x
, RECORDBYTES = .rambytes[vmode(x)], BATCHBYTES = getOption("ffbatchbytes"), ...)
## S3 method for class 'ffdf'
chunk(x
, RECORDBYTES = sum(.rambytes[vmode(x)]), BATCHBYTES = getOption("ffbatchbytes"), ...)
```

**Arguments**

x	ff or ffdf
RECORDBYTES	optional integer scalar representing the bytes needed to process an element of the ff_vector a single row of the ffdf
BATCHBYTES	integer scalar limiting the number of bytes to be processed in one chunk, default from <code>getOption("ffbatchbytes")</code> , see also <code>.rambytes</code>
...	further arguments passed to <code>chunk</code>

**Value**

A list with `ri` indexes each representing one chunk

**Author(s)**

Jens Oehlschlägel

**See Also**

`chunk`, `ffdf`

**Examples**

```
x <- data.frame(x=as.double(1:26), y=factor(letters), z=ordered(LETTERS))
a <- as.ffdf(x)
ceiling(26 / (300 %% sum(.rambytes[vmode(a)])))
chunk(a, BATCHBYTES=300)
ceiling(13 / (100 %% sum(.rambytes[vmode(a)])))
chunk(a, from=1, to = 13, BATCHBYTES=100)
rm(a); gc()

message("dummy example for linear regression with biglm on ffdf")
library(biglm)

message("NOTE that . in formula requires calculating terms manually
  because . as a data-dependant term is not allowed in biglm")
form <- Sepal.Length ~ Sepal.Width + Petal.Length + Petal.Width + Species

lmfit <- lm(form, data=iris)

firis <- as.ffdf(iris)
for (i in chunk(firis, by=50)){
  if (i[[1]]==1){
    message("first chunk is: ", i[[1]],":",i[[2]])
    biglmfit <- biglm(form, data=firis[i,,drop=FALSE])
  }else{
    message("next chunk is: ", i[[1]],":",i[[2]])
    biglmfit <- update(biglmfit, firis[i,,drop=FALSE])
  }
}
```

```
summary(lmfit)
summary(biglmfit)
stopifnot(all.equal(coef(lmfit), coef(biglmfit)))
```

---

clone

*Cloning ff and ram objects*


---

### Description

clone physically duplicates ff (and ram) objects and can additionally change some features, e.g. length.

### Usage

```
clone(x, ...)
## S3 method for class 'ff'
clone(x
, initdata = x
, length = NULL
, levels = NULL
, ordered = NULL
, dim = NULL
, dimorder = NULL
, bydim = NULL
, symmetric = NULL
, fixdiag = NULL
, names = NULL
, dimnames = NULL
, ramclass = NULL
, ramattrs = NULL
, vmode = NULL
, update = NULL
, pattern = NULL
, filename = NULL
, overwrite = FALSE
, pagesize = NULL
, caching = NULL
, finalizer = NULL
, finonexit = NULL
, FF_RETURN = NULL
, BATCHSIZE = .Machine$integer.max
, BATCHBYTES = getOption("ffbatchbytes")
, VERBOSE = FALSE
, ...)
## S3 method for class 'list'
clone(x, ...)
## Default S3 method:
clone(x, ...)
```



**Arguments**

x	x
initdata	scalar or vector of the <a href="#">.vimplemented vmodes</a> , recycled if needed, default 0, see also <a href="#">as.vmode</a> and <a href="#">vector.vmode</a>
length	optional vector <a href="#">length</a> of the object (default: derive from 'initdata' or 'dim'), see <a href="#">length.ff</a>
levels	optional character vector of levels if (in this case initdata must be composed of these) (default: derive from initdata)
ordered	indicate whether the levels are ordered (TRUE) or non-ordered factor (FALSE, default)
dim	optional array <a href="#">dim</a> , see <a href="#">dim.ff</a> and <a href="#">array</a>
dimorder	physical layout (default 1:length(dim)), see <a href="#">dimorder</a> and <a href="#">aperm</a>
bydim	dimorder by which to interpret the 'initdata', generalization of the 'byrow' paramter in <a href="#">matrix</a>
symmetric	extended feature: TRUE creates symmetric matrix (default FALSE)
fixdiag	extended feature: non-NULL scalar requires fixed diagonal for symmetric matrix (default NULL is free diagonal)
names	see <a href="#">names</a>
dimnames	NOT taken from initdata, see <a href="#">dimnames</a>
ramclass	class attribute attached when moving all or parts of this ff into ram, see <a href="#">ramclass</a>
ramattribs	additional attributes attached when moving all or parts of this ff into ram, see <a href="#">ramattribs</a>
vmode	virtual storage mode (default: derive from 'initdata'), see <a href="#">vmode</a> and <a href="#">as.vmode</a>
update	set to FALSE to avoid updating with 'initdata' (default TRUE) (used by <a href="#">ffdf</a> )
pattern	root pattern for automatic ff filename creation (default "ff"), see also <a href="#">physical</a>
filename	ff <a href="#">filename</a> (default tmpfile with 'pattern' prefix), see also <a href="#">physical</a>
overwrite	set to TRUE to allow overwriting existing files (default FALSE)
pagesize	pagesize in bytes for the memory mapping (default from <a href="#">getOptions("ffpagesize")</a> initialized by <a href="#">getdefaultpagesize</a> ), see also <a href="#">physical</a>
caching	caching scheme for the backend, currently 'mmnoflush' or 'mmeachflush' (flush mmpages at each swap, default from <a href="#">getOptions("ffcaching")</a> initialized with 'memorymap'), see also <a href="#">physical</a>
finalizer	name of finalizer function called when ff object is <a href="#">removed</a> , (default "deleteIfOpen" from <a href="#">getOptions("fffinalizer")</a> ), standard finalizers are <a href="#">close.ff</a> , <a href="#">delete.ff</a> and <a href="#">deleteIfOpen.ff</a> , see also <a href="#">reg.finalizer</a>
finonexit	logical scalar determining whether finalizer is also called when R is closed via <a href="#">q</a> , (default TRUE from <a href="#">getOptions("fffionexit")</a> )
FF_RETURN	logical scalar or ff object to be used. The default NULL creates a ff or ram clone, TRUE returns a ff clone, FALSE returns a ram clone. Handing over an ff object here uses this or stops if not <a href="#">ffsuitable</a>

BATCHSIZE	integer scalar limiting the number of elements to be processed in <code>update.ff</code> when <code>length(initdata)&gt;1</code> , default from <code>getOption("ffbatchsize")</code>
BATCHBYTES	integer scalar limiting the number of bytes to be processed in <code>update.ff</code> when <code>length(initdata)&gt;1</code> , default from <code>getOption("ffbatchbytes")</code> , see also <code>.rambytes</code>
VERBOSE	set to TRUE for verbosing in <code>update.ff</code> when <code>length(initdata)&gt;1</code> , default FALSE
...	further arguments to the generic

### Details

`clone` is generic. `clone.ff` is the workhorse behind `as.ram` and `as.ff`. For creating the desired object it calls `ff` which calls `update` for initialization.

### Value

an `ff` or `ram` object

### Author(s)

Jens Oehlschlägel

### See Also

`ff`, `update`, `as.ram`, `as.ff`

### Examples

```
x <- ff(letters, levels=letters)
y <- clone(x, length=52)
rm(x,y); gc()
```

---

clone.ffdf

*Cloning ffdf objects*

---

### Description

`clone` physically duplicates `ffdf` objects

### Usage

```
## S3 method for class 'ffdf'
clone(x, nrow=NULL, ...)
```

### Arguments

<code>x</code>	an <code>ffdf</code>
<code>nrow</code>	optionally the desired number of rows in the new object. Currently this works only together with <code>initdata=NULL</code>
...	further arguments passed to <code>clone</code> (usually not usefull)

**Details**

Creates a deep copy of an ffdF object by cloning all [physical](#) components including the [row.names](#)

**Value**

An object of type [ffdf](#)

**Author(s)**

Jens Oehlschlägel

**See Also**

[clone](#), [ffdf](#)

**Examples**

```
x <- as.ffdf(data.frame(a=1:26, b=letters))

message("Here we change the content of both x and y by reference")
y <- x
x$a[1] <- -1
y$a[1]

message("Here we change the content only of x because y is a deep copy")
y <- clone(x)
x$a[2] <- -2
y$a[2]
rm(x, y); gc()
```

---

close.ff

*Closing ff files*

---

**Description**

Close frees the Memory Mapping resources and closes the ff file without deleting the file data.

**Usage**

```
## S3 method for class 'ff'
close(con, ...)
## S3 method for class 'ffdf'
close(con, ...)
## S3 method for class 'ff_pointer'
close(con, ...)
```

**Arguments**

con            an open ff object  
 ...            ...

**Details**

The `ff_pointer` method is not intended for manual use, it is used at finalizer dispatch time. Closing `ffdf` objects will close all of their `physical` components including their `row.names` if they are `is.ff`

**Value**

TRUE if the file could be closed, FALSE if it was closed already (or NA if not all components of an `ffdf` returned FALSE or TRUE on closing)

**Author(s)**

Jens Oehlschlägel

**See Also**

[ff](#), [open.ff](#), [delete](#), [deleteIfOpen](#)

**Examples**

```
x <- ff(1:12)
close(x)
x
open(x)
x
rm(x); gc()
```

---

delete

*Deleting the file behind an ff object*

---

**Description**

The generic `delete` deletes the content of an object without removing the object itself. The generic `deleteIfOpen` does the same, but only if `is.open` returns TRUE.

**Usage**

```
delete(x, ...)
deleteIfOpen(x, ...)
## S3 method for class 'ff'
delete(x, ...)
## S3 method for class 'ffdf'
delete(x, ...)
## S3 method for class 'ff_pointer'
```

```

delete(x, ...)
## Default S3 method:
delete(x, ...)
## S3 method for class 'ff'
deleteIfOpen(x, ...)
## S3 method for class 'ff_pointer'
deleteIfOpen(x, ...)

```

### Arguments

x                    an ff or ram object  
...                    further arguments (not used)

### Details

The proper sequence to fully delete an ff object is: `delete(x);rm(x)`, where `delete.ff` frees the Memory Mapping resources and deletes the ff file, leaving intact the R-side object including its `class`, `physical` and `virtual` attributes. The default method is a compatibility function doing something similar with ram objects: by assigning an empty list to the name of the ram object to the parent frame we destroy the content of the object, leaving an empty stub that prevents raising an error if the parent frame calls the `delete(x);rm(x)` sequence.

The `deleteIfOpen` does the same as `delete` but protects closed ff objects from deletion, it is mainly intended for use through a finalizer, as are the `ff_pointer` methods.

### Value

`delete` returns TRUE if the/all ff files could be removed and FALSE otherwise.

`deleteIfOpen` returns TRUE if the/all ff files could be removed, FALSE if not and NA if the ff object was open.

### Note

Deletion of ff files can be triggered automatically via three routes:

1. if an ff object with a 'delete' finalizer is removed
2. if an ff object was created with `fffinonexit=TRUE` the finalizer is also called when R shuts down.
3. if an ff object was created in `getOption("fftempdir")`, it will be unlinked together with the `fftempdir` [.onUnload](#)

Thus in order to retain an ff file, one has to create it elsewhere than in `fftempdir` with a finalizer that does not destroy the file (by default files outside `fftempdir` get a 'close' finalizer) i.e. one of the following:

1. name the file AND use `fffinalizer="close"`
2. name the file AND use `fffinalizer="deleteIfOpen"` AND close the ff object before leaving R
3. name the file AND use `fffinalizer="delete"` AND use `fffinonexit=FALSE`

**Author(s)**

Jens Oehlschlägel

**See Also**[ff](#), [close.ff](#), [open.ff](#), [reg.finalizer](#)**Examples**

```
message('create the ff file outside getOption("fftempir"),
  it will have default finalizer "close", so you need to delete it explicately')
x <- ff(1:12, pattern="./ffexample")
delete(x)
rm(x)
```

dim.ff

*Getting and setting dim and dimorder***Description**

Assigning `dim` to an `ff_vector` changes it to an `ff_array`. Beyond that `dimorder` can be assigned to change from column-major order to row-major order or generalizations for higher order `ff_array`.

**Usage**

```
## S3 method for class 'ff'
dim(x)
## S3 method for class 'ffdf'
dim(x)
## S3 replacement method for class 'ff'
dim(x) <- value
## S3 replacement method for class 'ffdf'
dim(x) <- value
  dimorder(x, ...)
  dimorder(x, ...) <- value
## Default S3 method:
dimorder(x, ...)
## S3 method for class 'ff_array'
dimorder(x, ...)
## S3 method for class 'ffdf'
dimorder(x, ...)
## S3 replacement method for class 'ff_array'
dimorder(x, ...) <- value
## S3 replacement method for class 'ffdf'
dimorder(x, ...) <- value # just here to catch forbidden assignments
```

**Arguments**

x	a ff object
value	an appropriate integer vector
...	further arguments (not used)

**Details**

dim and dimorder are [virtual](#) attributes. Thus two copies of an R ff object can point to the same file but interpret it differently. dim has the usual meaning, dimorder defines the dimension order of storage, i.e. c(1,2) corresponds to R's standard column-major order, c(1,2) corresponds to row-major order, and for higher dimensional arrays dimorder can also be used. Standard dimorder is 1:length(dim(x)).

For [ffdf](#) dim returns the number of rows and virtual columns. With dim<- .ffdf only the number of rows can be changed. For convenience you can assign NA to the number of columns.

For [ffdf](#) the dimorder returns non-standard dimorder if any of its columns contains a ff object with non-standard dimorder (see [dimorderStandard](#)) An even higher level of virtualization is available using virtual windows, see [vw](#).

**Value**

names returns a character vector (or NULL)

**Note**

x[] returns a matrix like x[,] and thus respects dimorder, while x[i:j] returns a vector and simply returns elements in the stored order. Check the corresponding example twice, in order to make sure you understand that for non-standard dimorder x[1:length(x)] is *not the same* as as.vector(x[]).

**Author(s)**

Jens Oehlschlägel

**See Also**

[dim](#), [dimnames.ff\\_array](#), [dimorderStandard](#), [vw](#), [virtual](#)

**Examples**

```
x <- ff(1:12, dim=c(3,4), dimorder=c(2:1))
y <- x
dim(y) <- c(4,3)
dimorder(y) <- c(1:2)
x
y
x[]
y[]
x[,bydim=c(2,1)]
y[,bydim=c(2,1)]
```

```

message("NOTE that x[] like x[,] returns a matrix (respects dimorder),")
message("while x[1:12] returns a vector IN STORAGE ORDER")
message("check the following examples twice to make sure you understand this")
x[,]
x[]
as.vector(x[])
x[1:12]
rm(x,y); gc()

## Not run:
message("some performance comparison between different dimorders")
n <- 100
m <- 100000
a <- ff(1L,dim=c(n,m))
b <- ff(1L,dim=c(n,m), dimorder=2:1)
system.time(lapply(1:n, function(i)sum(a[i,])))
system.time(lapply(1:n, function(i)sum(b[i,])))
system.time(lapply(1:n, function(i){i<-(i-1)*(m/n)+1; sum(a[,i:(i+m/n-1)]))})
system.time(lapply(1:n, function(i){i<-(i-1)*(m/n)+1; sum(b[,i:(i+m/n-1)]))})
rm(a,b); gc()

## End(Not run)

```

---

dimnames.ffdf

*Getting and setting dimnames of ffdf*


---

## Description

Getting and setting dimnames, columnnames or rownames

## Usage

```

## S3 method for class 'ffdf'
dimnames(x)
## S3 replacement method for class 'ffdf'
dimnames(x) <- value
## S3 method for class 'ffdf'
names(x)
## S3 replacement method for class 'ffdf'
names(x) <- value
## S3 method for class 'ffdf'
row.names(x)
## S3 replacement method for class 'ffdf'
row.names(x) <- value

```

## Arguments

x a *ffdf* object  
value a character vector, or, for dimnames a list with two character vectors



**Details**

It is recommended not to assign row.names to a large ffd object.

**Value**

The assignment function return the changed ffd object. The other functions return the expected.

**Author(s)**

Jens Oehlschlägel

**See Also**

[ffdf](#), [dimnames.ff](#), [rownames](#), [colnames](#)

**Examples**

```
ffdf <- as.ffdf(data.frame(a=1:26, b=letters))
dimnames(ffdf)
row.names(ffdf) <- letters
dimnames(ffdf)
ffdf
rm(ffdf); gc()
```

---

dimnames.ff\_array      *Getting and setting dimnames*

---

**Description**

For ff\_arrays you can set dimnames.

**Usage**

```
## S3 method for class 'ff_array'
dimnames(x)
## S3 replacement method for class 'ff_array'
dimnames(x) <- value
```

**Arguments**

x	a ff array (or matrix)
value	a list with length(dim(x)) elements (either NULL or character vector of length of dimension)

**Details**

if [vw](#) is set, [dimnames.ff\\\_array](#) returns the appropriate part of the names, but you can't set dimnames while [vw](#) is set. [dimnames](#) returns NULL for ff\_vectors and setting dimnames for ff\_vector is not allowed, but setting [names](#) is.

**Value**

dimnames returns a list, see [dimnames](#)

**Author(s)**

Jens Oehlschlägel

**See Also**

[dimnames](#), [dim.ff](#), [names.ff](#), [vw](#), [virtual](#)

**Examples**

```
x <- ff(1:12, dim=c(3,4), dimnames=list(letters[1:3], LETTERS[1:4]))
dimnames(x)
dimnames(x) <- list(LETTERS[1:3], letters[1:4])
dimnames(x)
dimnames(x) <- NULL
dimnames(x)
rm(x); gc()
```

---

dimorderCompatible      *Test for dimorder compatibility*

---

**Description**

dimorderStandard returns TRUE if the dimorder is standard (ascending), vectorStandard returns TRUE if the dimorder-bydim combination is compatible with a standard elementwise vector interpretation, dimorderCompatible returns TRUE if two dimorders have a compatible elementwise vector interpretation and vectorCompatible returns TRUE if dimorder-bydim combinations have a compatible elementwise vector interpretation.

**Usage**

```
dimorderStandard(dimorder)
vectorStandard(dimorder, bydim = NULL)
dimorderCompatible(dim, dim2, dimorder, dimorder2)
vectorCompatible(dim, dim2, dimorder=NULL, dimorder2=NULL, bydim = NULL, bydim2 = NULL)
```

**Arguments**

dim	a <a href="#">dim</a>
dim2	a <a href="#">dim</a>
dimorder	a <a href="#">dimorder</a>
dimorder2	a <a href="#">dimorder</a>
bydim	a bydim order, see <a href="#">[.ff]</a>
bydim2	a bydim order, see argument fromdim in <a href="#">update.ff</a>

**Value**

TRUE if compatibility has been detected, FALSE otherwise

**Note**

does not yet gurantee to detect all compatible configurations, but the most important ones

**Author(s)**

Jens Oehlschlägel

**See Also**

[dimorder](#), [ffconform](#)

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<code>dummy.dimnames</code>	<i>Array: make dimnames</i>
-----------------------------	-----------------------------

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**Description**

makes standard dimnames from letters and integers (for testing)

**Usage**

```
dummy.dimnames(x)
```

**Arguments**

x                    an [array](#)

**Value**

a list with character vectors suitable to be assigned as dimnames to x

**Author(s)**

Jens Oehlschlägel

**See Also**

[dimnames](#)

**Examples**

```
dummy.dimnames(matrix(1:12, 3, 4))
```

## Description

These are the main methods for reading and writing data from ff files.

## Usage

```
## S3 method for class 'ff'
x[i, pack = FALSE]
## S3 replacement method for class 'ff'
x[i, add = FALSE, pack = FALSE] <- value
## S3 method for class 'ff_array'
x[..., bydim = NULL, drop = getOption("ffdrop"), pack = FALSE]
## S3 replacement method for class 'ff_array'
x[..., bydim = NULL, add = FALSE, pack = FALSE] <- value
## S3 method for class 'ff'
x[[i]]
## S3 replacement method for class 'ff'
x[[i, add = FALSE]] <- value
```

## Arguments

x	an ff object
i	missing OR a single index expression OR a <a href="#">hi</a> object
...	missing OR up to length(dim(x)) index expressions OR <a href="#">hi</a> objects
drop	logical scalar indicating whether array dimensions shall be dropped
bydim	the dimorder which shall be used in interpreting vector to/from array data
pack	FALSE to prevent rle-packing in hybrid index preprocessing, see <a href="#">as.hi</a>
value	the values to be assigned, possibly recycled
add	TRUE if the values should rather increment than overwrite at the target positions, see <a href="#">readwrite.ff</a>

## Details

The single square bracket operators `[` and `[<-` are the workhorses for accessing the content of an ff object. They support `ff_vector` and `ff_array` access ([dim.ff](#)), they respect virtual windows ([vw](#)), [names.ff](#) and [dimnames.ff](#) and retain [ramclass](#) and [ramattribs](#) and thus support `POSIXct` and `factor`, see [levels.ff](#).

The functionality of `[` and `[<-` can be combined into one efficient operation, see [swap](#).

The double square bracket operator `[[` is a shortcut for [get.ff](#) resp. [set.ff](#), however, you should not rely on this for the future, see [LimWarn](#). For programming please prefer `[`.

**Value**

The read operators `[` and `[[` return data from the `ff` object, possibly decorated with `names`, `dim`, `dimnames` and further attributes and classes (see `ramclass`, `ramattribs`)

The write operators `[<-` and `[[<-` return the 'modified' `ff` object (like all assignment operators do).

**Index expressions**

```
x <- ff(1:12, dim=c(3,4), dimnames=list(letters[1:3], NULL))
```

<i>allowed expression</i>	–	example
positive integers		<code>x[ 1 ,1]</code>
negative integers		<code>x[ -(2:12) ]</code>
logical		<code>x[ c(TRUE, FALSE, FALSE) ,1]</code>
character		<code>x[ "a" ,1]</code>
integer matrices		<code>x[ rbind(c(1,1)) ]</code>
hybrid index		<code>x[ hi ,1]</code>
<i>disallowed expression</i>	–	example
zeros		<code>x[ 0 ]</code>
NAs		<code>x[ NA ]</code>

**Dimorder and bydim**

Arrays in R have always standard `dimorder` `1:length(dim(x))` while `ff` allows to store an array in a different dimorder. Using nonstandard dimorder (see `dimorderStandard`) can speed up certain access operations: while `matrix dimorder=c(1,2)` – column-major order – allows fast extraction of columns, `dimorder=c(2,1)` allows fast extraction of rows.

While the `dimorder` – being an attribute of an `ff_array` – controls how the vector in an `ff` file is interpreted, the `bydim` argument to the extractor functions controls, how assignment vector values in `[<-` are translated to the array and how the array is translated to a vector in `[` subscripting. Note that `bydim=c(2,1)` corresponds to `matrix(..., byrow=TRUE)`.

**Multiple vector interpretation in arrays**

In case of non-standard `dimorder` (see `dimorderStandard`) the vector sequence of array elements in R and in the `ff` file differs. To access array elements in file order, you can use `getset.ff`, `readwrite.ff` or copy the `ff` object and set `dim(ff)<-NULL` to get a vector view into the `ff` object (using `[` dispatches the vector method `[.ff`). To access the array elements in R standard `dimorder` you simply use `[` which dispatches to `[.ff_array`. Note that in this case `as.hi` will unpack the complete index, see next section.

**RAM expansion of index expressions**

Some index expressions do not consume RAM due to the `hi` representation, for example `1:n` will almost consume no RAM however large `n`. However, some index expressions are expanded and require to `maxindex(i) * .rambytes["integer"]` bytes, either because the sorted sequence of index positions cannot be rle-packed efficiently or because `hiparse` cannot yet parse such expression and falls back to evaluating/expanding the index expression. If the index positions are not sorted,

the index will be expanded and a second vector is needed to store the information for re-ordering, thus the index requires  $2 * \text{maxindex}(i) * \text{.rambytes["integer"]}$  bytes.

### RAM expansion when recycling assignment values

Some assignment expressions do not consume RAM for recycling, for example `x[1:n] <- 1:k` will not consume RAM however large `n` compared to `k`, when `x` has standard `dimorder`. However, if `length(value)>1`, assignment expressions with non-ascending index positions trigger recycling the value R-side to the full index length. This will happen if `dimorder` does not match parameter `bydim` or if the index is not sorted ascending.

### Author(s)

Jens Oehlschlägel

### See Also

[ff](#), [swap](#), [add](#), [readwrite.ff](#), [LimWarn](#)

### Examples

```
message("look at different dimorders")
x <- ff(1:12, dim=c(3,4), dimorder=c(1,2))
x[]
as.vector(x[])
x[1:12]
x <- ff(1:12, dim=c(3,4), dimorder=c(2,1))
x[]
as.vector(x[])
message("Beware (might be changed)")
x[1:12]

message("look at different bydim")
matrix(1:12, nrow=3, ncol=4, byrow=FALSE)
x <- ff(1:12, dim=c(3,4), bydim=c(1,2))
x
matrix(1:12, nrow=3, ncol=4, byrow=TRUE)
x <- ff(1:12, dim=c(3,4), bydim=c(2,1))
x
x[, bydim=c(2,1)]
as.vector(x[, bydim=c(2,1)])
message("even consistent interpretation of vectors in assignments")
x[, bydim=c(1,2)] <- x[, bydim=c(1,2)]
x
x[, bydim=c(2,1)] <- x[, bydim=c(2,1)]
x
rm(x); gc()

## Not run:
message("some performance implications of different dimorders")
n <- 100
m <- 100000
```

```

a <- ff(1L,dim=c(n,m))
b <- ff(1L,dim=c(n,m), dimorder=2:1)
system.time(lapply(1:n, function(i)sum(a[i,])))
system.time(lapply(1:n, function(i)sum(b[i,])))
system.time(lapply(1:n, function(i){i<-(i-1)*(m/n)+1; sum(a[,i:(i+m/n-1)])}))
system.time(lapply(1:n, function(i){i<-(i-1)*(m/n)+1; sum(b[,i:(i+m/n-1)])}))

n <- 100
a <- ff(1L,dim=c(n,n,n))
b <- ff(1L,dim=c(n,n,n), dimorder=4:1)
system.time(lapply(1:n, function(i)sum(a[i,,,])))
system.time(lapply(1:n, function(i)sum(a[,i,,])))
system.time(lapply(1:n, function(i)sum(a[,,,i])))
system.time(lapply(1:n, function(i)sum(a[,,,i])))
system.time(lapply(1:n, function(i)sum(b[i,,,])))
system.time(lapply(1:n, function(i)sum(b[,i,,])))
system.time(lapply(1:n, function(i)sum(b[,,,i])))
system.time(lapply(1:n, function(i)sum(b[,,,i])))

n <- 100
m <- 100000
a <- ff(1L,dim=c(n,m))
b <- ff(1L,dim=c(n,m), dimorder=2:1)
system.time(ffrowapply(sum(a[i1:i2,]), a, RETURN=TRUE, CFUN="csum", BATCHBYTES=16104816%/%20))
system.time(ffcolapply(sum(a[,i1:i2]), a, RETURN=TRUE, CFUN="csum", BATCHBYTES=16104816%/%20))
system.time(ffrowapply(sum(b[i1:i2,]), b, RETURN=TRUE, CFUN="csum", BATCHBYTES=16104816%/%20))
system.time(ffcolapply(sum(b[,i1:i2]), b, RETURN=TRUE, CFUN="csum", BATCHBYTES=16104816%/%20))

rm(a,b); gc()

## End(Not run)

```

---

Extract.ffdf

*Reading and writing data.frames (ffdf)*


---

## Description

These are the main methods for reading and writing data from `ffdf` objects.

## Usage

```

## S3 method for class 'ffdf'
x[i, j, drop = ncols == 1]
## S3 replacement method for class 'ffdf'
x[i, j] <- value
## S3 method for class 'ffdf'
x[[i, j, exact = TRUE]]
## S3 replacement method for class 'ffdf'
x[[i, j]] <- value

```

```
## S3 method for class 'ffdf'
x$i
## S3 replacement method for class 'ffdf'
x$i <- value
```

### Arguments

x	an ff object
i	a row subscript or a matrix subscript or a list subscript
j	a column subscript
drop	logical. If TRUE the result is coerced to the lowest possible dimension. The default is to drop if only one column is left, but not to drop if only one row is left.
value	A suitable replacement value: it will be repeated a whole number of times if necessary and it may be coerced: see the Coercion section. If NULL, deletes the column if a single column is selected with <code>[[&lt;-</code> or <code>\$&lt;-</code> .
exact	logical: see <code>[</code> , and applies to column names.

### Details

The subscript methods `[`, `[[` and `$`, behave symmetrical to the assignment functions `<-`, `[[<-` and `$<-`. What the former return is the assignment value to the latter. A notable exception is assigning NULL in `[[<-` and `$<-` which removes the [virtual](#) column from the ffdf (and the [physical](#) component if it is no longer needed by any virtual column). Creating new columns via `[[<-` and `$<-` requires giving a name to the new column (character subscripting). `<-` does not allow to create new columns, only to replace existing ones.

### Subscript expressions and return values

<i>allowed expression</i>	– example	– returnvalue
row selection	<code>x[i, ]</code>	<a href="#">data.frame</a> or single row as list if <code>drop=TRUE</code> , like from <code>data.frame</code>
column selection	<code>x[, i]</code>	<a href="#">data.frame</a> or single column as vector unless <code>drop=TRUE</code> , like from <code>data.frame</code>
matrix selection	<code>x[cbind(i, j)]</code>	vector of the integer-matrix indexed cells (if the column types are compatible)
virtual selection	<code>x[i]</code>	<a href="#">ffdf</a> with the selected columns only
physical selection	<code>x[[i]]</code>	the selected <a href="#">ff</a>
physical selection	<code>x\$i</code>	the selected <a href="#">ff</a>

### Author(s)

Jens Oehlschlägel

### See Also

[ffdf](#), [Extract.data.frame](#), [Extract.ff](#)



**Examples**

```

d <- data.frame(a=letters, b=rev(letters), c=1:26)
x <- as.ffdf(d)

d[, ]
x[, ]

d[1:2, ]
x[1:2, ]

d[, 1]
x[, 1]

d[, 1:2]
x[, 1:2]

d[cbind(1:2, 2:1)]
x[cbind(1:2, 2:1)]

d[1]
x[1]

d[[1]]
x[[1]]

d$a
x$a

d$a[1:2]
x$a[1:2]

rm(x); gc()

```

ff

*ff classes for representing (large) atomic data***Description**

The ff package provides atomic data structures that are stored on disk but behave (almost) as if they were in RAM by mapping only a section (pagesize) into main memory (the effective main memory consumption per ff object). Several access optimization techniques such as Hybrid Index Preprocessing ([as.hi](#), [update.ff](#)) and Virtualization ([virtual](#), [vt](#), [vw](#)) are implemented to achieve good performance even with large datasets. In addition to the basic access functions, the ff package also provides compatibility functions that facilitate writing code for ff and ram objects ([clone](#), [as.ff](#), [as.ram](#)) and very basic support for operating on ff objects ([ffapply](#)). While the (possibly packed) raw data is stored on a flat file, meta informations about the atomic data structure such as its dimension, virtual storage mode ([vmode](#)), factor level encoding, internal length etc.. are stored as an ordinary R object (external pointer plus attributes) and can be saved in the workspace. The raw flat file data encoding is always in native machine format for optimal performance and provides

several packing schemes for different data types such as logical, raw, integer and double (in an extended version support for more tightly packed virtual data types is supported). flatfile data files can be shared among ff objects in the same R process or even from different R processes due to Memory-Mapping, although the caching effects have not been tested extensively.

Please do read and understand the limitations and warnings in [LimWarn](#) before you do anything serious with package ff.

## Usage

```
ff( initdata = NULL
  , length   = NULL
  , levels   = NULL
  , ordered  = NULL
  , dim      = NULL
  , dimorder = NULL
  , bydim    = NULL
  , symmetric = FALSE
  , fixdiag  = NULL
  , names    = NULL
  , dimnames = NULL
  , ramclass = NULL
  , ramattribs = NULL
  , vmode    = NULL
  , update   = NULL
  , pattern  = NULL
  , filename = NULL
  , overwrite = FALSE
  , readonly  = FALSE
  , pagesize  = NULL # getOption("ffpagesize")
  , caching   = NULL # getOption("ffcaching")
  , finalizer = NULL
  , finonexit = NULL # getOption("fffinonexit")
  , FF_RETURN = TRUE
  , BATCHSIZE = .Machine$integer.max
  , BATCHBYTES = getOption("ffbatchbytes")
  , VERBOSE   = FALSE
)
```

## Arguments

initdata	scalar or vector of the <a href="#">.vimplemented vmodes</a> , recycled if needed, default 0, see also <a href="#">as.vmode</a> and <a href="#">vector.vmode</a>
length	optional vector <a href="#">length</a> of the object (default: derive from 'initdata' or 'dim'), see <a href="#">length.ff</a>
levels	optional character vector of levels if (in this case initdata must be composed of these) (default: derive from initdata)
ordered	indicate whether the levels are ordered (TRUE) or non-ordered factor (FALSE, default)

dim	optional array <a href="#">dim</a> , see <a href="#">dim.ff</a> and <a href="#">array</a>
dimorder	physical layout (default 1:length(dim)), see <a href="#">dimorder</a> and <a href="#">aperm</a>
bydim	dimorder by which to interpret the 'initdata', generalization of the 'byrow' paramter in <a href="#">matrix</a>
symmetric	extended feature: TRUE creates symmetric matrix (default FALSE)
fixdiag	extended feature: non-NULL scalar requires fixed diagonal for symmetric matrix (default NULL is free diagonal)
names	NOT taken from initdata, see <a href="#">names</a>
dimnames	NOT taken from initdata, see <a href="#">dimnames</a>
ramclass	class attribute attached when moving all or parts of this ff into ram, see <a href="#">ramclass</a>
ramattribs	additional attributes attached when moving all or parts of this ff into ram, see <a href="#">ramattribs</a>
vmode	virtual storage mode (default: derive from 'initdata'), see <a href="#">vmode</a> and <a href="#">as.vmode</a>
update	set to FALSE to avoid updating with 'initdata' (default TRUE) (used by <a href="#">ffdf</a> )
pattern	root pattern with or without path for automatic ff filename creation (default NULL translates to "ff"), see also argument 'filename'
filename	ff <a href="#">filename</a> with or without path (default tmpfile with 'pattern' prefix); without path the file is created in <code>getOption("fftmpdir")</code> , with path '.' the file is created in <a href="#">getwd</a> . Note that files created in <code>getOption("fftmpdir")</code> have default finalizer "delete" while other files have default finalizer "close". See also arguments 'pattern' and 'finalizer' and <a href="#">physical</a>
overwrite	set to TRUE to allow overwriting existing files (default FALSE)
readonly	set to TRUE to forbid writing to existing files
pagesize	pagesize in bytes for the memory mapping (default from <code>getOption("ffpagesize")</code> ) initialized by <a href="#">getdefaultpagesize</a> , see also <a href="#">physical</a>
caching	caching scheme for the backend, currently 'mmnoflush' or 'mmeachflush' (flush mmpages at each swap, default from <code>getOption("ffcaching")</code> ) initialized with 'mmeachflush', see also <a href="#">physical</a>
finalizer	name of finalizer function called when ff object is <a href="#">removed</a> (default: ff files created in <code>getOption("fftmpdir")</code> are considered temporary and have default finalizer <a href="#">delete</a> , files created in other locations have default finalizer <a href="#">close</a> ); available finalizer generics are "close", "delete" and "deleteIfOpen", available methods are <a href="#">close.ff</a> , <a href="#">delete.ff</a> and <a href="#">deleteIfOpen.ff</a> , see also argument 'finonexit' and <a href="#">finalizer</a>
finonexit	logical scalar determining whether and <a href="#">finalize</a> is also called when R is closed via <a href="#">q</a> , (default TRUE from <code>getOption("fffiononexit")</code> )
FF_RETURN	logical scalar or ff object to be used. The default TRUE creates a new ff file. FALSE returns a ram object. Handing over an ff object here uses this or stops if not <a href="#">ffsuitable</a>
BATCHSIZE	integer scalar limiting the number of elements to be processed in <a href="#">update.ff</a> when <code>length(initdata)&gt;1</code> , default from <code>.Machine\$integer.max</code>
BATCHBYTES	integer scalar limiting the number of bytes to be processed in <a href="#">update.ff</a> when <code>length(initdata)&gt;1</code> , default from <code>getOption("ffbatchbytes")</code> , see also <a href="#">rambytes</a>
VERBOSE	set to TRUE for verbosing in <a href="#">update.ff</a> when <code>length(initdata)&gt;1</code> , default FALSE

## Details

The atomic data is stored in `filename` as a native encoded raw flat file on disk, OS specific limitations of the file system apply. The number of elements per ff object is limited to the integer indexing, i.e. `.Machine$integer.max`. Atomic objects created with ff are `is.open`, a C++ object is ready to access the file via memory-mapping. Currently the C++ backend provides two caching schemes: `'mmnoflush'` let the OS decide when to flash memory mapped pages and `'mmeachflush'` will flush memory mapped pages at each page swap per ff file. These minimal memory resources can be released by `close`ing or `delete`ing the ff file. ff objects can be `saved` and `loaded` across R sessions. If the ff file still exists in the same location, it will be `opened` automatically at the first attempt to access its data. If the ff object is `removed`, at the next garbage collection (see `gc`) the ff object's `finalizer` is invoked. Raw data files can be made accessible as an ff object by explicitly given the filename and vmode but no size information (length or dim). The ff object will open the file and handle the data with respect to the given vmode. The `close` finalizer will close the ff file, the `delete` finalizer will delete the ff file. The default finalizer `deleteIfOpen` will delete open files and do nothing for closed files. If the default finalizer is used, two actions are needed to protect the ff file against deletion: create the file outside the standard `'fftempdir'` and close the ff object before removing it or before quitting R. When R is exited through `q`, the finalizer will be invoked depending on the `'ffinonexit'` option, furthermore the `'fftempdir'` is `unlinked`.

## Value

If (`!FF_RETURN`) then a ram object like those generated by `vector`, `matrix`, `array` but with attributes `'vmode'`, `'physical'` and `'virtual'` accessible via `vmode`, `physical` and `virtual`

If (`FF_RETURN`) an object of class `'ff'` which is a list with two components:

<code>physical</code>	an external pointer of class <code>'ff_pointer'</code> which carries attributes with copy by reference semantics: changing a physical attribute of a copy changes the original
<code>virtual</code>	an empty list which carries attributes with copy by value semantics: changing a virtual attribute of a copy does not change the original

## Physical object component

The `'ff_pointer'` carries the following `'physical'` or `readonly` attributes, which are accessible via `physical`:

<code>vmode</code>	see <code>vmode</code>
<code>maxlength</code>	see <code>maxlength</code>
<code>pattern</code>	see parameter <code>'pattern'</code>
<code>filename</code>	see <code>filename</code>
<code>pagesize</code>	see parameter <code>'pagesize'</code>
<code>caching</code>	see parameter <code>'caching'</code>
<code>finalizer</code>	see parameter <code>'finalizer'</code>
<code>finonexit</code>	see parameter <code>'finonexit'</code>
<code>readonly</code>	see <code>is.readonly</code>
<code>class</code>	The external pointer needs class <code>'ff_pointer'</code> to allow method dispatch of finalizers

## Virtual object component

The 'virtual' component carries the following attributes (some of which might be NULL):

Length	see <a href="#">length.ff</a>
Levels	see <a href="#">levels.ff</a>
Names	see <a href="#">names.ff</a>
VW	see <a href="#">vw.ff</a>
Dim	see <a href="#">dim.ff</a>
Dimorder	see <a href="#">dimorder</a>
Symmetric	see <a href="#">symmetric.ff</a>
Fixdiag	see <a href="#">fixdiag.ff</a>
ramclass	see <a href="#">ramclass</a>
ramattribs	see <a href="#">ramattribs</a>

## Class

You should not rely on the internal structure of ff objects or their ram versions. Instead use the accessor functions like [vmode](#), [physical](#) and [virtual](#). Still it would be wise to avoid attributes AND classes 'vmode', 'physical' and 'virtual' in any other packages. Note that the 'ff' object's class attribute also has copy-by-value semantics ('virtual'). For the 'ff' object the following class attributes are known:

vector	c("ff_vector", "ff")
matrix	c("ff_matrix", "ff_array", "ff")
array	c("ff_array", "ff")
symmetric matrix	c("ff_symm", "ff")
distance matrix	c("ff_dist", "ff_symm", "ff")
reserved for future use	c("ff_mixed", "ff")

## Methods

The following methods and functions are available for ff objects:

<i>Type</i>	<i>Name</i>	<i>Assign</i>	<i>Comment</i>
			<b>Basic functions</b>
function	<a href="#">ff</a>		constructor for ff and ram objects
generic	<a href="#">update</a>		updates one ff object with the content of another
generic	<a href="#">clone</a>		clones an ff object optionally changing some of its features
method	<a href="#">print</a>		print ff
method	<a href="#">str</a>		ff object structure
			<b>Class test and coercion</b>
function	<a href="#">is.ff</a>		check if inherits from ff
generic	<a href="#">as.ff</a>		coerce to ff, if not yet
generic	<a href="#">as.ram</a>		coerce to ram retaining some of the ff information
generic	<a href="#">as.bit</a>		coerce to <a href="#">bit</a>
			<b>Virtual storage mode</b>
generic	<a href="#">vmode</a>	<-	get and set virtual mode (setting only for ram, not for ff objects)

generic	<code>as.vmode</code>		coerce to vmode (only for ram, not for ff objects)
			<b>Physical attributes</b>
function	<code>physical</code>	<-	set and get physical attributes
generic	<code>filename</code>	<-	get and set filename
generic	<code>pattern</code>	<-	get pattern and set filename path and prefix via pattern
generic	<code>maxlength</code>		get maxlength
generic	<code>is.sorted</code>	<-	set and get if is marked as sorted
generic	<code>na.count</code>	<-	set and get NA count, if set to non-NA only swap methods can change and na.count is
generic	<code>is.readonly</code>		get if is readonly
			<b>Virtual attributes</b>
function	<code>virtual</code>	<-	set and get virtual attributes
method	<code>length</code>	<-	set and get length
method	<code>dim</code>	<-	set and get dim
generic	<code>dimorder</code>	<-	set and get the order of dimension interpretation
generic	<code>vt</code>		virtually transpose ff_array
method	<code>t</code>		create transposed clone of ff_array
generic	<code>vw</code>	<-	set and get virtual windows
method	<code>names</code>	<-	set and get names
method	<code>dimnames</code>	<-	set and get dimnames
generic	<code>symmetric</code>		get if is symmetric
generic	<code>fixdiag</code>	<-	set and get fixed diagonal of symmetric matrix
method	<code>levels</code>	<-	levels of factor
generic	<code>recodelevels</code>		recode a factor to different levels
generic	<code>sortlevels</code>		sort the levels and recoce a factor
method	<code>is.factor</code>		if is factor
method	<code>is.ordered</code>		if is ordered (factor)
generic	<code>ramclass</code>		get ramclass
generic	<code>ramattribs</code>		get ramattribs
			<b>Access functions</b>
function	<code>get.ff</code>		get single ff element (currently <code>[[</code> is a shortcut)
function	<code>set.ff</code>		set single ff element (currently <code>[[&lt;-</code> is a shortcut)
function	<code>getset.ff</code>		set single ff element and get old value in one access operation
function	<code>read.ff</code>		get vector of contiguous elements
function	<code>write.ff</code>		set vector of contiguous elements
function	<code>readwrite.ff</code>		set vector of contiguous elements and get old values in one access operation
method	<code>[</code>		get vector of indexed elements, uses HIP, see <a href="#">hi</a>
method	<code>[&lt;-</code>		set vector of indexed elements, uses HIP, see <a href="#">hi</a>
generic	<code>swap</code>		set vector of indexed elements and get old values in one access operation
generic	<code>add</code>		(almost) unifies '+=' operation for ff and ram objects
generic	<code>bigsample</code>		sample from ff object
			<b>Opening/Closing/Deleting</b>
generic	<code>is.open</code>		check if ff is open
method	<code>open</code>		open ff object (is done automatically on access)
method	<code>close</code>		close ff object (releases C++ memory and protects against file deletion if <code>deleteIfOpen</code> )
generic	<code>delete</code>		deletes ff file (unconditionally)
generic	<code>deleteIfOpen</code>		deletes ff file if ff object is open (finalization method)
generic	<code>finalizer</code>	<-	get and set finalizer
generic	<code>finalize</code>		force finalization

		<b>Other</b>
function	<a href="#">geterror.ff</a>	get error code
function	<a href="#">geterrstr.ff</a>	get error message

## ff options

Through [options](#) or [getOption](#) one can change and query global features of the ff package:

<i>option</i>	<i>description</i>	<i>default</i>
fftempdir	default directory for creating ff files	<a href="#">tempdir</a>
fffinalizer	name of default finalizer	<a href="#">deleteIfOpen</a>
fffinonexit	default for invoking finalizer on exit of R	TRUE
ffpagesize	default pagesize	<a href="#">getdefaultpagesize</a>
ffcaching	caching scheme for the C++ backend	'mmnoflush'
ffdrops	default for the 'drop' parameter in the ff subscript methods	TRUE
ffbatchbytes	default for the byte limit in batched/chunked processing	<a href="#">memory.limit()</a> %% 100

## OS specific

The following table gives an overview of file size limits for common file systems (see [http://en.wikipedia.org/wiki/Comparison\\_of\\_file\\_systems](http://en.wikipedia.org/wiki/Comparison_of_file_systems) for further details):

<b>File System</b>	<b>File size limit</b>
FAT16	2GB
FAT32	4GB
NTFS	16GB
ext2/3/4	16GB to 2TB
ReiserFS	4GB (up to version 3.4) / 8TB (from version 3.5)
XFS	8EB
JFS	4PB
HFS	2GB
HFS Plus	16GB
USF1	4GB to 256TB
USF2	512GB to 32PB
UDF	16EB

## Credits

Package Version 1.0

Daniel Adler	<dadler@uni-goettingen.de> R package design, C++ generic file vectors, Memory-Mapping, 64-bit Multi-Indexing adapter and Document
Oleg Nenadic	<onenadi@uni-goettingen.de> Index sequence packing, Documentation
Walter Zucchini	<wzucchi@uni-goettingen.de> Array Indexing, Sampling, Documentation

Christian Gläser <christian\\_glaeser@gmx.de>  
 Wrapper for biglm package

#### Package Version 2.0

Jens Oehlschlägel <Jens.Oehlschlaegel@truecluster.com>

R package redesign; Hybrid Index Preprocessing; transparent object creation and finalization; vmode design

Daniel Adler <dadler@uni-goettingen.de>

C++ generic file vectors, vmode implementation and low-level bit-packing/unpacking, arithmetic operations

### Licence

Package under GPL-2, included C++ code released by Daniel Adler under the less restrictive ISCL

### Note

Note that the standard finalizers are generic functions, their dispatch to the 'ff\_pointer' method happens at finalization time, their 'ff' methods exist for direct calling.

### See Also

[vector](#), [matrix](#), [array](#), [as.ff](#), [as.ram](#)

### Examples

```
message("make sure you understand the following ff options
before you start using the ff package!!")
oldoptions <- options(fffinalizer="deleteIfOpen", fffinonexit="TRUE", fftempdir=tempdir())
message("an integer vector")
ff(1:12)
message("a double vector of length 12")
ff(0, 12)
message("a 2-bit logical vector of length 12 (vmode='boolean' has 1 bit)")
ff(vmode="logical", length=12)
message("an integer matrix 3x4 (standard colwise physical layout)")
ff(1:12, dim=c(3,4))
message("an integer matrix 3x4 (rowwise physical layout, but filled in standard colwise order)")
ff(1:12, dim=c(3,4), dimorder=c(2,1))
message("an integer matrix 3x4 (standard colwise physical layout, but filled in rowwise order
aka matrix(, byrow=TRUE))")
ff(1:12, dim=c(3,4), bydim=c(2,1))
gc()
options(oldoptions)

if (ffxtensions()){
  message("a 26-dimensional boolean array using 1-bit representation
(file size 8 MB compared to 256 MB int in ram)")
  a <- ff(vmode="boolean", dim=rep(2, 26))
  dimnames(a) <- dummy.dimnames(a)
}
```



```

    rm(a); gc()
  }

## Not run:

message("This 2GB biglm example can take long, you might want to change
  the size in order to define a size appropriate for your computer")
require(biglm)

b <- 1000
n <- 100000
k <- 3
memory.size(max = TRUE)
system.time(
x <- ff(vmode="double", dim=c(b*n,k), dimnames=list(NULL, LETTERS[1:k]))
)
memory.size(max = TRUE)
system.time(
ffrowapply({
  l <- i2 - i1 + 1
  z <- rnorm(l)
  x[i1:i2,] <- z + matrix(rnorm(l*k), l, k)
}, X=x, VERBOSE=TRUE, BATCHSIZE=n)
)
memory.size(max = TRUE)

form <- A ~ B + C
first <- TRUE
system.time(
ffrowapply({
  if (first){
    first <- FALSE
    fit <- biglm(form, as.data.frame(x[i1:i2,],drop=FALSE))
  }else
    fit <- update(fit, as.data.frame(x[i1:i2,],drop=FALSE))
}, X=x, VERBOSE=TRUE, BATCHSIZE=n)
)
memory.size(max = TRUE)
first
fit
summary(fit)
rm(x); gc()

## End(Not run)

```

## Description

The `ffapply` functions support convenient batched processing of `ff` objects such that each single batch or chunk will not exhaust RAM and such that batches have sizes as similar as possible, see [bbatch](#). Differing from R's standard `apply` which applies a `FUN`ction, the `ffapply` functions do apply an `EXPR`ession and provide two indices `FROM="i1"` and `TO="i2"`, which mark beginning and end of the batch and can be used in the applied expression. Note that the `ffapply` functions change the two indices in their parent frame, to avoid conflicts you can use different names through `FROM="i1"` and `TO="i2"`. For support of creating return values see details.

## Usage

```
ffvecapply(EXPR, X = NULL, N = NULL, VMODE = NULL, VBYTES = NULL, RETURN = FALSE
, CFUN = NULL, USE.NAMES = TRUE, FF_RETURN = TRUE, BREAK = ".break"
, FROM = "i1", TO = "i2"
, BATCHSIZE = .Machine$integer.max, BATCHBYTES = getOption("ffbatchbytes")
, VERBOSE = FALSE)
ffrowapply(EXPR, X = NULL, N = NULL, NCOL = NULL, VMODE = NULL, VBYTES = NULL
, RETURN = FALSE, RETCOL = NCOL, CFUN = NULL, USE.NAMES = TRUE, FF_RETURN = TRUE
, FROM = "i1", TO = "i2"
, BATCHSIZE = .Machine$integer.max, BATCHBYTES = getOption("ffbatchbytes")
, VERBOSE = FALSE)
ffcolapply(EXPR, X = NULL, N = NULL, NROW = NULL, VMODE = NULL, VBYTES = NULL
, RETURN = FALSE, RETROW = NROW, CFUN = NULL, USE.NAMES = TRUE, FF_RETURN = TRUE
, FROM = "i1", TO = "i2"
, BATCHSIZE = .Machine$integer.max, BATCHBYTES = getOption("ffbatchbytes")
, VERBOSE = FALSE)
ffapply(EXPR = NULL, AFUN = NULL, MARGIN = NULL, X = NULL, N = NULL, DIM = NULL
, VMODE = NULL, VBYTES = NULL, RETURN = FALSE, CFUN = NULL, USE.NAMES = TRUE
, FF_RETURN = TRUE, IDIM = "idim"
, FROM = "i1", TO = "i2", BREAK = ".break"
, BATCHSIZE = .Machine$integer.max, BATCHBYTES = getOption("ffbatchbytes")
, VERBOSE = FALSE)
```

## Arguments

<code>EXPR</code>	the <a href="#">expression</a> to be applied
<code>AFUN</code>	<code>ffapply</code> only: alternatively to <code>EXPR</code> the name of a function to be applied, automatically converted to <code>EXPR</code>
<code>MARGIN</code>	<code>ffapply</code> only: the margins along which to loop in <code>ffapply</code>
<code>X</code>	an <code>ff</code> object from which several parameters can be derived, if they are not given directly: <code>N</code> , <code>NCOL</code> , <code>NROW</code> , <code>DIM</code> , <code>VMODE</code> , <code>VBYTES</code> , <code>FF_RETURN</code>
<code>N</code>	the total number of elements in the loop, e.g. number of elements in <code>ffvecapply</code> or number of rows in <code>ffrowapply</code>
<code>NCOL</code>	<code>ffrowapply</code> only: the number of columns needed to calculate batch sizes
<code>NROW</code>	<code>ffcolapply</code> only: the number of rows needed to calculate batch sizes
<code>DIM</code>	<code>ffapply</code> only: the dimension of the array needed to calculate batch sizes

VMODE	the <a href="#">vmode</a> needed to prepare the RETURN object and to derive VBYTES if they are not given directly
VBYTES	the bytes per cell – see <a href="#">.rambytes</a> – to calculate the RAM requirements per cell
BATCHBYTES	the max number of bytes per batch, default <code>getOption("ffbatchbytes")</code>
BATCHSIZE	an additional restriction on the number of loop elements, default <code>=.Machine\$integer.max</code>
FROM	the name of the index that marks the beginning of the batch, default 'i1', change if needed to avoid naming-conflicts in the calling frame
TO	the name of the index that marks the end of the batch, default 'i2', change if needed to avoid naming-conflicts in the calling frame
IDIM	ffapply only: the name of an R variable used for loop-switching, change if needed to avoid naming-conflicts in the calling frame
BREAK	ffapply only: the name of an R object in the calling frame that triggers break out of the batch loop, if 1) it exists 2) is.logical and 3) is TRUE
RETURN	TRUE to prepare a return value (default FALSE)
CFUN	name of a collapsing function, see <a href="#">CFUN</a>
RETCOL	NULL gives return vector[1:N], RETCOL gives return matrix[1:N, 1:RETCOL]
RETROW	NULL gives return vector[1:N], RETROW gives return matrix[1:RETROW, 1:N]
FF_RETURN	FALSE to return a ram object, TRUE to return an ff object, or an ff object that is <a href="#">ffsuitable</a> to absorb the return data
USE.NAMES	FALSE to suppress attaching names or dimnames to the result
VERBOSE	TRUE to verbose the batches

## Details

`ffvecapply` is the simplest `ffapply` method for `ff_vectors`. `ffrowapply` and `ffcolapply` is for `ff_matrix`, and `ffapply` is the most general method for `ff_arrays` and `ff_vectors`.

There are many ways to change the return value of the `ffapply` functions. In its simplest usage – batched looping over an expression – they don't return anything, see [invisible](#). If you switch `RETURN=TRUE` in `ffvecapply` then it is assumed that all looped expressions together return one vector of length N, and via parameter `FF_RETURN`, you can decide whether this vector is in ram or is an ff object (or even which ff object to use). `ffrowapply` and `ffcolapply` additionally have parameter `RETCOL` resp. `RETROW` which defaults to returning a matrix of the original size; in order to just return a vector of length N set this to NULL, or specify a number of columns/rows for the return matrix. It is assumed that the expression will return appropriate pieces for this return structure (see examples). If you specify `RETURN=TRUE` and a collapsing function name `CFUN`, then it is assumed that the batched expressions return aggregated information, which is first collected in a list, and finally the collapsing function is called on this list: `do.call(CFUN, list)`. If you want to return the unmodified list, you have to specify `CFUN="list"` for obvious reasons.

`ffapply` allows usages not completely unlike [apply](#): you can specify the name of a function `AFUN` to be applied over `MARGIN`. However note that you must specify `RETURN=TRUE` in order to get a return value. Also note that currently `ffapply` assumes that your expression returns exactly one value per cell in `DIM[MARGINS]`. If you want to return something more complicated, you MUST specify a `CFUN="list"` and your return value will be a list with dim attribute `DIM[MARGINS]`. This means that for a function `AFUN` returning a scalar, `ffapply` behaves very similar to [apply](#), see examples.

Note also that `ffapply` might create a object named `'ffapply.dimexhausted'` in its parent frame, and it uses a variable in the parent frame for loop-switching between dimensions, the default name `'idim'` can be changed using the `IDIM` parameter. Finally you can break out of the implied loops by assigning `TRUE` to a variable with the name in `BREAK`.

### Value

see details

### Note

xx The complete generation of the return value is preliminary and the arguments related to defining the return value might still change, especially `ffapply` is work in progress

### Author(s)

Jens Oehlschlägel

### See Also

[apply](#), [expression](#), [bbatch](#), [repfromto](#), [ffsuitable](#)

### Examples

```
message("ffvecapply examples")
x <- ff(vmode="integer", length=1000)
message("loop evaluate expression without returning anything")
ffvecapply(x[i1:i2] <- i1:i2, X=x, VERBOSE=TRUE)
ffvecapply(x[i1:i2] <- i1:i2, X=x, BATCHSIZE=200, VERBOSE=TRUE)
ffvecapply(x[i1:i2] <- i1:i2, X=x, BATCHSIZE=199, VERBOSE=TRUE)
message("lets return the combined expressions as a new ff object")
ffvecapply(i1:i2, N=length(x), VMODE="integer", RETURN=TRUE, BATCHSIZE=200)
message("lets return the combined expressions as a new ram object")
ffvecapply(i1:i2, N=length(x), VMODE="integer", RETURN=TRUE, FF_RETURN=FALSE, BATCHSIZE=200)
message("lets return the combined expressions in existing ff object x")
x[] <- 0L
ffvecapply(i1:i2, N=length(x), VMODE="integer", RETURN=TRUE, FF_RETURN=x, BATCHSIZE=200)
x
message("aggregate and collapse")
ffvecapply(summary(x[i1:i2]), X=x, RETURN=TRUE, CFUN="list", BATCHSIZE=200)
ffvecapply(summary(x[i1:i2]), X=x, RETURN=TRUE, CFUN="rbind", BATCHSIZE=200)
ffvecapply(summary(x[i1:i2]), X=x, RETURN=TRUE, CFUN="cmean", BATCHSIZE=200)

message("how to do colSums with ffrowapply")
x <- ff(1:10000, vmode="integer", dim=c(1000, 10))
ffrowapply(colSums(x[i1:i2],drop=FALSE), X=x, RETURN=TRUE, CFUN="list", BATCHSIZE=200)
ffrowapply(colSums(x[i1:i2],drop=FALSE), X=x, RETURN=TRUE, CFUN="rbind", BATCHSIZE=200)
ffrowapply(colSums(x[i1:i2],drop=FALSE), X=x, RETURN=TRUE, CFUN="csum", BATCHSIZE=200)

message("further ffrowapply examples")
x <- ff(1:10000, vmode="integer", dim=c(1000, 10))
message("loop evaluate expression without returning anything")
```

```

ffrowapply(x[i1:i2, ] <- i1:i2, X=x, BATCHSIZE=200)
message("lets return the combined expressions as a new ff object (x unchanged)")
ffrowapply(2*x[i1:i2, ], X=x, RETURN=TRUE, BATCHSIZE=200)
message("lets return a single row aggregate")
ffrowapply(t(apply(x[i1:i2,,drop=FALSE], 1, mean)), X=x, RETURN=TRUE, RETCOL=NULL, BATCHSIZE=200)
message("lets return a 6 column aggregates")
y <- ffrowapply( t(apply(x[i1:i2,,drop=FALSE], 1, summary)), X=x
, RETURN=TRUE, RETCOL=length(summary(0)), BATCHSIZE=200)
colnames(y) <- names(summary(0))
y
message("determine column minima if a complete column does not fit into RAM")
ffrowapply(apply(x[i1:i2,], 2, min), X=x, RETURN=TRUE, CFUN="pmin", BATCHSIZE=200)

message("ffapply examples")
x <- ff(1:720, dim=c(8,9,10))
dimnames(x) <- dummy.dimnames(x)
message("apply function with scalar return value")
apply(X=x[,], MARGIN=3:2, FUN=sum)
apply(X=x[,], MARGIN=2:3, FUN=sum)
ffapply(X=x, MARGIN=3:2, AFUN="sum", RETURN=TRUE, BATCHSIZE=8)
message("this is what CFUN is based on")
ffapply(X=x, MARGIN=2:3, AFUN="sum", RETURN=TRUE, CFUN="list", BATCHSIZE=8)

message("apply functions with vector or array return value currently have limited support")
apply(X=x[,], MARGIN=3:2, FUN=summary)
message("you must use CFUN, the rest is up to you")
y <- ffapply(X=x, MARGIN=3:2, AFUN="summary", RETURN=TRUE, CFUN="list", BATCHSIZE=8)
y
y[[1]]

rm(x); gc()

```

ffconform

*Get most conforming argument***Description**

ffconform returns position of 'most' conformable ff argument or zero if the arguments are not conforming

**Usage**

```
ffconform(..., vmode = NULL, fail = "stop")
```

**Arguments**

...	two or more ff objects
vmode	handing over target vmode here supresses searching for a common vmode, see <a href="#">maxffmode</a>
fail	the name of a function to call if not-conforming, default <a href="#">stop</a>

## Details

A reference argument is defined to be the first argument with a `dim` attribute or the longest vector. The other arguments are then compared to the reference to check for conformity, which is violated if `vmodes` are not conforming or if the reference has not a multiple length of each other or if the dimensions do not match or if we have a `dimorder` conflict because not all arguments have the same `dimorderStandard`.

## Value

the position of the most conforming argument or 0 (zero) if not conforming.

## Note

xx Work in progress for package **R.ff**

## Author(s)

Jens Oehlschlägel

## See Also

[ffsuitable](#), [maxffmode](#), [ymismatch](#), [stop](#), [warning](#), [dimorderStandard](#)

## Examples

```
a <- ff(1:10)
b <- clone(a)
c <- ff(1:20)
d <- ff(1:21)
ffconform(a,b)
ffconform(c,a)
ffconform(a,c)
ffconform(c,a,b)

d1 <- ff(1:20, dim=c(2,10))
d2 <- ff(1:20, dim=c(10,2))
ffconform(c,d1)
ffconform(c,d2)
ffconform(d1,c)
ffconform(d2,c)
try(ffconform(d1,d2))
ffconform(d1,d1)

rm(a,b,c,d1,d2); gc()
```

ffdf

*ff class for data.frames***Description**

Function 'ffdf' creates ff data.frames stored on disk very similar to 'data.frame'

**Usage**

```
ffdf(...
, row.names = NULL
, ff_split = NULL
, ff_join = NULL
, ff_args = NULL
, update = TRUE
, BATCHSIZE = .Machine$integer.max
, BATCHBYTES = getOption("ffbatchbytes")
, VERBOSE = FALSE)
```

**Arguments**

...	ff vectors or matrices (optionally wrapped in I()) that shall be bound together to an ffdf object
row.names	A <a href="#">character</a> vector. Not recommended for large objects with many rows.
ff_split	A vector of character names or integer positions identifying input components to physically split into single ff_vectors. If vector elements have names, these are used as root name for the new ff files.
ff_join	A list of vectors with character names or integer positions identifying input components to physically join in the same ff matrix. If list elements have names, these are used to name the new ff files.
update	By default (TRUE) new ff files are updated with content of input ff objects. Setting to FALSE prevents this update.
ff_args	a list with further arguments passed to ff in case that new ff objects are created via 'ff_split' or 'ff_join'
BATCHSIZE	passed to <a href="#">update.ff</a>
BATCHBYTES	passed to <a href="#">update.ff</a>
VERBOSE	passed to <a href="#">update.ff</a>

**Details**

By default, creating an 'ffdf' object will NOT create new ff files, instead existing files are referenced. This differs from [data.frame](#), which always creates copies of the input objects, most notably in `data.frame(matrix())`, where an input matrix is converted to single columns. ffdf by contrast, will store an input matrix physically as the same matrix and virtually map it to columns. Physically copying a large ff matrix to single ff vectors can be expensive. More generally, ffdf

objects have a [physical](#) and a [virtual](#) component, which allows very flexible dataframe designs: a physically stored matrix can be virtually mapped to single columns, a couple of physically stored vectors can be virtually mapped to a single matrix. The means to configure these are `I` for the virtual representation and the `'ff_split'` and `'ff_join'` arguments for the physical representation. An ff matrix wrapped into `'I()'` will return the input matrix as a single object, using `'ff_split'` will store this matrix as single vectors - and thus create new ff files. `'ff_join'` will copy a couple of input vectors into a unified new ff matrix with `dimorder=c(2,1)`, but virtually they will remain single columns. The returned ffd object has also a `dimorder` attribute, which indicates whether the ffd object contains a matrix with non-standard dimorder `c(2,1)`, see [dimorderStandard](#). Currently, [virtual windows](#) are not supported for ffd.

## Value

A list with components

<code>physical</code>	the underlying ff vectors and matrices, to be accessed via <a href="#">physical</a>
<code>virtual</code>	the virtual features of the ffd including the virtual-to-physical mapping, to be accessed via <a href="#">virtual</a>
<code>row.names</code>	the optional row.names, see argument row.names

and class `'ffdf'` (NOTE that ffd does not inherit from ff)

## Methods

The following methods and functions are available for ffd objects:

<i>Type</i>	<i>Name</i>	<i>Assign</i>	<i>Comment</i>
<b>Basic functions</b>			
function	<a href="#">ffdf</a>		constructor for ffd objects
generic	<a href="#">update</a>		updates one ffd object with the content of another
generic	<a href="#">clone</a>		clones an ffd object
method	<a href="#">print</a>		print ffd
method	<a href="#">str</a>		ffdf object structure
<b>Class test and coercion</b>			
function	<a href="#">is.ffdf</a>		check if inherits from ff
generic	<a href="#">as.ffdf</a>		coerce to ff, if not yet
generic	<a href="#">as.data.frame</a>		coerce to ram data.frame
<b>Virtual storage mode</b>			
generic	<a href="#">vmode</a>		get virtual modes for all (virtual) columns
<b>Physical attributes</b>			
function	<a href="#">physical</a>		get physical attributes
<b>Virtual attributes</b>			
function	<a href="#">virtual</a>		get virtual attributes
method	<a href="#">length</a>		get length
method	<a href="#">dim</a>	<-	get dim and set nrow
generic	<a href="#">dimorder</a>		get the dimorder (non-standard if any component is non-standard)
method	<a href="#">names</a>	<-	set and get names
method	<a href="#">row.names</a>	<-	set and get row.names
method	<a href="#">dimnames</a>	<-	set and get dimnames



method	<code>pattern</code>	<-	set pattern (rename/move files)
			<b>Access functions</b>
method	<code>[</code>	<-	set and get data.frame content ( <code>[, ]</code> ) or get ffd with less columns ( <code>[[]</code> )
method	<code>[[</code>	<-	set and get single column ff object
method	<code>\$</code>	<-	set and get single column ff object
			<b>Opening/Closing/Deleting</b>
generic	<code>is.open</code>		tri-bool is.open status of the physical ff components
method	<code>open</code>		open all physical ff objects (is done automatically on access)
method	<code>close</code>		close all physical ff objects
method	<code>delete</code>		deletes all physical ff files
method	<code>finalize</code>		call finalizer
			<b>processing</b>
method	<code>chunk</code>		create chunked index
method	<code>sortLevels</code>		sort and recode levels
			<b>Other</b>

**Note**

Note that in theory, accessing a chunk of rows from a matrix with `dimorder=c(2,1)` should be faster than accessing across a bunch of vectors. However, at least under windows, the OS has difficulties filecaching parts from very large files, therefore - until we have partitioning - the recommended physical storage is in single vectors.

**Author(s)**

Jens Oehlschlägel

**See Also**

[data.frame](#), [ff](#), for more example see [physical](#)

**Examples**

```
m <- matrix(1:12, 3, 4, dimnames=list(c("r1","r2","r3"), c("m1","m2","m3","m4")))
v <- 1:3
ffm <- as.ff(m)
ffv <- as.ff(v)
```

```
d <- data.frame(m, v)
ffd <- ffd(fm, v=ffv, row.names=row.names(ffm))
all.equal(d, ffd[,])
ffd
physical(ffd)
```

```
d <- data.frame(m, v)
ffd <- ffd(fm, v=ffv, row.names=row.names(ffm), ff_split=1)
all.equal(d, ffd[,])
ffd
physical(ffd)
```

```

d <- data.frame(m, v)
ffdf <- ffdf(ffm, v=ffv, row.names=row.names(ffm), ff_join=list(newff=c(1,2)))
all.equal(d, ffdf[,])
ffdf
physical(ffdf)

d <- data.frame(I(m), I(v))
ffdf <- ffdf(m=I(ffm), v=I(ffv), row.names=row.names(ffm))
all.equal(d, ffdf[,])
ffdf
physical(ffdf)

rm(ffm,ffv,ffdf); gc()

```

ffdfindexget

*Reading and writing ffdf data.frame using ff subscripts***Description**

Function `ffdfindexget` allows to extract rows from an `ffdf` data.frame according to positive integer subscripts stored in an `ff` vector.

Function `ffdfindexset` allows the inverse operation: assigning to rows of an `ffdf` data.frame according to positive integer subscripts stored in an `ff` vector. These functions allow more control than the method dispatch of `[]` and `[]<-` if an `ff` integer subscript is used.

**Usage**

```

ffdfindexget(x, index, indexorder = NULL, autoindexorder = 3, FF_RETURN = NULL
, BATCHSIZE = NULL, BATCHBYTES = getOption("ffmaxbytes"), VERBOSE = FALSE)
ffdfindexset(x, index, value, indexorder = NULL, autoindexorder = 3
, BATCHSIZE = NULL, BATCHBYTES = getOption("ffmaxbytes"), VERBOSE = FALSE)

```

**Arguments**

<code>x</code>	A <code>ffdf</code> data.frame containing the elements
<code>index</code>	A <code>ff</code> integer vector with integer subscripts in the range from 1 to <code>length(x)</code> .
<code>value</code>	A <code>ffdf</code> data.frame like <code>x</code> with the rows to be assigned
<code>indexorder</code>	Optionally the return value of <code>ffindexorder</code> , see details
<code>autoindexorder</code>	The minimum number of columns (which need chunked indexing) for which we switch from on-the-fly ordering to stored <code>ffindexorder</code>
<code>FF_RETURN</code>	Optionally an <code>ffdf</code> data.frame of the same type as <code>x</code> in which the returned values shall be stored, see details.
<code>BATCHSIZE</code>	Optimal limit for the batchsize (see details)
<code>BATCHBYTES</code>	Limit for the number of bytes per batch
<code>VERBOSE</code>	Logical scalar for verbosing

## Details

Accessing rows of an `ffdf` data.frame identified by integer positions in an `ff` vector is a non-trivial task, because it could easily lead to random-access to disk files. We avoid random access by loading batches of the subscript values into RAM, order them ascending, and only then access the `ff` values on disk. Such ordering is done on-the-fly for upto `autoindexorder-1` columns that need ordering. For `autoindexorder` or more columns we do the batched ordering upfront with `ffindexorder` and then re-use it in each call to `ffindexget` resp. `ffindexset`.

## Value

Function `ffdfindexget` returns a `ffdf` data.frame with those rows selected by the `ff` index vector. Function `ffdfindexset` returns `x` with those rows replaced that had been requested by `index` and `value`.

## Author(s)

Jens Oehlschlägel

## See Also

[Extract.ff](#), [ffindexget](#), [ffindexorder](#)

## Examples

```
message("ff integer subscripts with ffdf return/assign values")
x <- ff(factor(letters))
y <- ff(1:26)
d <- ffdf(x,y)
i <- ff(2:9)
di <- d[i,]
di
d[i,] <- di
message("ff integer subscripts: more control with ffindexget/ffindexset")
di <- ffdfindexget(d, i, FF_RETURN=di)
d <- ffdfindexset(d, i, di)
rm(x, y, d, i, di)
gc()
```

## Description

These functions allow convenient sorting and ordering of collections of (`ff`) vectors organized in (`ffdf`) data.frames

## Usage

```
dforder(x, ...)  
dfsrt(x, ...)  
ramdforder(x, ...)  
ramdfsrt(x, ...)  
ffdforder(x, ...)  
ffdfsrt(x, ...)
```

## Arguments

`x` a [data.frame](#) (for `dforder`, `dfsrt`, `ramorder`, `ramsrt`) or an [ffdf](#) object (for `ffdforder`, `ffdfsrt`)

`...` further arguments passed to [sort](#), [ramsrt](#) or [ffsrt](#) (for objects with one column) or passed to [order](#), [ramorder](#) or [fforder](#) (for objects with multiple columns)

## Value

the order functions return an (ff) vector of integer order positions, the sort functions return a sorted clone of the (ffdf) input data.frame

## Author(s)

Jens Oehlschlägel

## See Also

[sort](#), [ramsrt](#) or [ffsrt](#)  
[order](#), [ramorder](#) or [fforder](#)

## Examples

```
x <- ff(sample(1e5, 1e6, TRUE))  
y <- ff(sample(1e5, 1e6, TRUE))  
z <- ff(sample(1e5, 1e6, TRUE))  
d <- ffdf(x, y, z)  
d2 <- ffdfsrt(d)  
d2  
d  
d2 <- d[1:2]  
i <- ffdforder(d2)  
d[i,]  
rm(x, y, z, i, d, d2)  
gc()
```

---

ffdrop	<i>Delete an ffarchive</i>
--------	----------------------------

---

**Description**

Delete the <file>.Rdata and <file>.ffData files behind an ffarchive

**Usage**

```
ffdrop(file)
```

**Arguments**

file                vector of archive filenames (without extensions)

**Value**

A list with components

RData                vector with results of [file.remove](#) on RData files

ffData                Description of 'comp2'

**Note**

This deletes file on disk without warning

**Author(s)**

Jens Oehlschlägel

**See Also**

[ffsave](#), [ffinfo](#), [ffload](#)

---

ffindexget	<i>Reading and writing ff vectors using ff subscripts</i>
------------	---

---

**Description**

Function `ffindexget` allows to extract elements from an ff vector according to positive integer subscripts stored in an ff vector.

Function `ffindexset` allows the inverse operation: assigning to elements of an ff vector according to positive integer subscripts stored in an ff vector. These functions allow more control than the method dispatch of `[]` and `[]<-` if an ff integer subscript is used.

**Usage**

```
ffindexget(x, index, indexorder = NULL, FF_RETURN = NULL
, BATCHSIZE = NULL, BATCHBYTES = getOption("ffmaxbytes"), VERBOSE = FALSE)
ffindexset(x, index, value, indexorder = NULL
, BATCHSIZE = NULL, BATCHBYTES = getOption("ffmaxbytes"), VERBOSE = FALSE)
```

**Arguments**

x	A <code>ff</code> vector containing the elements
index	A <code>ff</code> integer vector with integer subscripts in the range from 1 to <code>length(x)</code> .
value	An <code>ff</code> vector of the same <code>vmode</code> as <code>x</code> containing the values to be assigned
indexorder	Optionally the return value of <code>ffindexorder</code> , see details
FF_RETURN	Optionally an <code>ff</code> vector of the same <code>vmode</code> as <code>x</code> in which the returned values shall be stored, see details.
BATCHSIZE	Optimal limit for the batchsize (see details)
BATCHBYTES	Limit for the number of bytes per batch
VERBOSE	Logical scalar for verbosing

**Details**

Accessing integer positions in an `ff` vector is a non-trivial task, because it could easily lead to random-access to a disk file. We avoid random access by loading batches of the subscript values into RAM, order them ascending, and only then access the `ff` values on disk. Since ordering is expensive, it may pay to do the batched ordering once upfront and then re-use it with `ffindexorder`, similar to storing and using hybrid index information with [as.hi](#).

**Value**

Function `ffindexget` returns an `ff` vector with the extracted elements.  
Function `ffindexset` returns the `ff` vector in which we have updated values.

**Author(s)**

Jens Oehlschlägel

**See Also**

[Extract.ff](#), [ffdfindexget](#), [ffindexorder](#)

**Examples**

```
message("ff integer subscripts with ff return/assign values")
x <- ff(factor(letters))
i <- ff(2:9)
xi <- x[i]
xi
xi[] <- NA
xi
```

```
x[i] <- xi
x
message("ff integer subscripts: more control with ffindexget/ffindexset")
xi <- ffindexget(x, i, FF_RETURN=xi)
x <- ffindexset(x, i, xi)
rm(x, i, xi)
gc()
```

ffindexorder

*Sorting: chunked ordering of integer subscript positions***Description**

Function `ffindexorder` will calculate chunkwise the order positions to sort all positions in a chunk ascending.

Function `ffindexordersize` does the calculation of the chunksize for `ffindexorder`.

**Usage**

```
ffindexordersize(length, vmode, BATCHBYTES = getOption("ffmaxbytes"))
ffindexorder(index, BATCHSIZE, FF_RETURN = NULL, VERBOSE = FALSE)
```

**Arguments**

<code>index</code>	A <code>ff</code> integer vector with integer subscripts.
<code>BATCHSIZE</code>	Limit for the chunksize (see details)
<code>BATCHBYTES</code>	Limit for the number of bytes per batch
<code>FF_RETURN</code>	Optionally an <code>ff</code> integer vector in which the chunkwise order positions are stored.
<code>VERBOSE</code>	Logical scalar for activating verbosing.
<code>length</code>	Number of elements in the index
<code>vmode</code>	The <code>vmode</code> of the <code>ff</code> vector to which the index shall be applied with <code>ffindexget</code> or <code>ffindexset</code>

**Details**

Accessing integer positions in an `ff` vector is a non-trivial task, because it could easily lead to random-access to a disk file. We avoid random access by loading batches of the subscript values into RAM, order them ascending, and only then access the `ff` values on disk. Such an ordering can be done on-the-fly by `ffindexget` or it can be created upfront with `ffindexorder`, stored and re-used, similar to storing and using hybrid index information with [as.hi](#).

**Value**

Function `ffindexorder` returns an `ff` integer vector with an attribute `BATCHSIZE` (the chunksize finally used, not the one given with argument `BATCHSIZE`).

Function `ffindexordersize` returns a balanced batchsize as returned from `bbatch`.

**Author(s)**

Jens Oehlschlägel

**See Also**

[ffindexget](#), [as.hi](#), [bbatch](#)

**Examples**

```
x <- ff(sample(40))
message("fforder requires sorting")
i <- fforder(x)
message("applying this order i is done by ffindexget")
x[i]
message("applying this order i requires random access,
  therefore ffindexget does chunkwise sorting")
ffindexget(x, i)
message("if we want to apply the order i multiple times,
  we can do the chunkwise sorting once and store it")
s <- ffindexordersize(length(i), vmode(i), BATCHBYTES = 100)
o <- ffindexorder(i, s$b)
message("this is how the stored chunkwise sorting is used")
ffindexget(x, i, o)
message("")
rm(x,i,s,o)
gc()
```

---

ffinfo

*Inspect content of ff saves*

---

**Description**

Find out which objects and ff files are in a pair of files saved with [ffsave](#)

**Usage**

```
ffinfo(file)
```

**Arguments**

**file** a character string giving the name (without extension) of the .RData and .ffData files to load



**Value**

a list with components

RData	a list, one element for each object (named like the object): a character vector with the names of the ff files
ffData	a list, one element for each path (names like the path): a character vector with the names of the ff files
rootpath	the root path relative to which the files are stored in the .ffData zip

**Note**

For large files and the zip64 format use zip 3.0 and unzip 6.0 from <https://sourceforge.net/projects/infozip/>.

**Author(s)**

Jens Oehlschlägel

**See Also**

[ffsave](#), [ffload](#), [ffdrop](#)

---

ffload	<i>Reload ffSaved Datasets</i>
--------	--------------------------------

---

**Description**

Reload datasets written with the function `ffsave` or `ffsave.image`.

**Usage**

```
ffload(file, list = character(0L), envir = parent.frame()
, rootpath = NULL, overwrite = FALSE)
```

**Arguments**

file	a character string giving the name (without extension) of the .RData and .ffData files to load
list	An optional vector of names selecting those objects to be restored (default NULL restores all)
envir	the environment where the data should be loaded.
rootpath	an optional rootpath where to restore the ff files (default NULL restores in the original location)
overwrite	logical indicating whether possibly existing ff files shall be overwritten

**Details**

`ffinfo` can be used to inspect the contents an ffsaved pair of `.RData` and `.ffData` files. Argument `list` can then be used to restore only part of the ffsave.

**Value**

A character vector with the names of the restored ff files

**Note**

The ff files are not platform-independent with regard to byte order. For large files and the zip64 format use zip 3.0 and unzip 6.0 from <https://sourceforge.net/projects/infozip/>.

**Author(s)**

Jens Oehlschlägel

**See Also**

[load](#), [ffsave](#), [ffinfo](#), [ffdrop](#)

---

fforder

*Sorting: order from ff vectors*

---

**Description**

Returns order with regard to one or more ff vectors

**Usage**

```
fforder(...  
  , index = NULL  
  , use.index = NULL  
  , aux = NULL  
  , auxindex = NULL  
  , has.na = TRUE  
  , na.last = TRUE  
  , decreasing = FALSE  
  , BATCHBYTES = getOption("ffmaxbytes")  
  , VERBOSE = FALSE  
)
```

**Arguments**

...	one of more ff vectors which define the order
index	an optional ff integer vector used to store the order output
use.index	A boolean flag telling fforder whether to use the positions in 'index' as input. If you do this, it is your responsibility to assure legal positions - otherwise you risk a crash.
aux	An optional named list of ff vectors that can be used for temporary copying – the names of the list identify the <a href="#">vmodes</a> for which the respective ff vector is suitable.
auxindex	An optional ff integer vector for temporary storage of integer positions.
has.na	boolean scalar telling fforder whether the vector might contain NAs. <i>Note</i> that you risk a crash if there are unexpected NAs with <code>has.na=FALSE</code>
na.last	boolean scalar telling fforder whether to order NAs last or first. <i>Note</i> that 'boolean' means that there is no third option NA as in <a href="#">order</a>
decreasing	boolean scalar telling fforder whether to order increasing or decreasing
BATCHBYTES	maximum number of RAM bytes fforder should try not to exceed
VERBOSE	cat some info about the ordering

**Details**

fforder tries to order the vector in-RAM, if not possible it uses (a yet simple) out-of-memory algorithm. Like [ramorder](#) the in-RAM ordering method is chosen depending on context information.

**Value**

An ff vector with the positions that are required to sort the input as specified – with an attribute [na.count](#) with as many values as columns in ...

**Author(s)**

Jens Oehlschlägel

**See Also**

[ramorder](#), [ffsort](#), [ffdforder](#), [ffindexget](#)

**Examples**

```
x <- ff(sample(1e5, 1e6, TRUE))
y <- ff(sample(1e5, 1e6, TRUE))
d <- ffd(x, y)

i <- fforder(y)
y[i]
i <- fforder(x, index=i)
x[i]
d[i,]
```

```
i <- fforder(x, y)
d[i,]

i <- ffdforder(d)
d[i,]

rm(x, y, d, i)
gc()
```

---

ffreturn	<i>Return suitable ff object</i>
----------	----------------------------------

---

### Description

ffreturn returns FF\_RETURN if it is [ffsuitable](#) otherwise creates a suitable [ffsuitable](#) object

### Usage

```
ffreturn(FF_RETURN = NULL, FF_PROTO = NULL, FF_ATTR = NULL)
```

### Arguments

FF_RETURN	the object to be tested for suitability
FF_PROTO	the prototype object which FF_RETURN should match
FF_ATTR	a list of additional attributes dominating those from FF_PROTO

### Value

a suitable [ffsuitable](#) object

### Note

xx Work in progress for package **R.ff**

### Author(s)

Jens Oehlschlägel

### See Also

[ffconform](#), [ffsuitable](#)

---

ffsave	<i>Save R and ff objects</i>
--------	------------------------------

---

### Description

ffsave writes an external representation of R and ff objects to an ffarchive. The objects can be read back from the file at a later date by using the function [ffload](#).

### Usage

```
ffsave(...
, list = character(0L)
, file = stop("'file' must be specified")
, envir = parent.frame()
, rootpath = NULL
, add = FALSE

, move = FALSE
, compress = !move
, compression_level = 6
, precheck=TRUE
)
ffsave.image(file = stop("'file' must be specified"), safe = TRUE, ...)
```

### Arguments

...	For ffsave the names of the objects to be saved (as symbols or character strings), for ffsave.image further arguments passed to ffsave
list	A character vector containing the names of objects to be saved.
file	A name for the the ffarchive, i.e. the two files <file>.RData and <file>.ffData
envir	environment to search for objects to be saved.
add	logical indicating whether the objects shall be added to the ffarchive (in this case rootpath is taken from an existing archive)
move	logical indicating whether ff files shall be moved instead of copied into the <file>.ffData
compress	logical specifying whether saving to a named file is to use compression.
compression_level	compression level passed to zip, default 6
rootpath	optional path component that all <i>all</i> ff files share and that can be dropped/replaced when calling <a href="#">ffload</a>
precheck	logical: should the existence of the objects be checked before starting to save (and in particular before opening the file/connection)?
safe	logical. If TRUE, a temporary file is used for creating the saved workspace. The temporary file is renamed to <file>.ffData if the save succeeds. This preserves an existing workspace <file>.ffData if the save fails, but at the cost of using extra disk space during the save.

## Details

ffsave stores objects and ff files in an ffarchive named <file>: i.e. it saves all specified objects via [save](#) in a file named <file>.RData and saves all ff files related to these objects in a zipfile named <file>.ffData using an external zip utility.

By default files are stored relative to the `rootpath=""` and will be restored relative to `\code{"\"` (in its original location). By providing a partial path prefix via argument `rootpath` the files are stored relative to this `rootpath`. The `rootpath` is stored in the <file>.RData with the name `.ff.rootpath`. I.e. even if the ff objects were saved with argument `rootpath` to `ffsave`, `ffload` by default restores in the original location. By using argument `rootpath` to `ffload` you can restore relative to a different `rootpath` (and using argument `rootpath` to `ffsave` gave you shorter relative paths)

By using argument `add` in `ffsave` you can add more objects to an existing ffarchive and by using argument `list` in `ffload` you can selectively restore objects.

The content of the ffarchive can be inspected using [ffinfo](#) before actually loading any of the objects.

The ffarchive can be deleted from disk using [ffdrop](#).

## Value

a character vector with messages returned from the zip utility (one for each ff file zipped)

## Note

The ff files are not platform-independent with regard to byte order. For large files and the zip64 format use `zip 3.0` and `unzip 6.0` from <https://sourceforge.net/projects/infozip/>.

## Author(s)

Jens Oehlschlägel

## See Also

[ffinfo](#) for inspecting the content of the ffarchive

[ffload](#) for loading all or some of the ffarchive

[ffdrop](#) for deleting one or more ffarchives

## Examples

```
## Not run:
message("let's create some ff objects")
n <- 8e3
a <- ff(sample(n, n, TRUE), vmode="integer", length=n, filename="d:/tmp/a.ff")
b <- ff(sample(255, n, TRUE), vmode="ubyte", length=n, filename="d:/tmp/b.ff")
x <- ff(sample(255, n, TRUE), vmode="ubyte", length=n, filename="d:/tmp/x.ff")
y <- ff(sample(255, n, TRUE), vmode="ubyte", length=n, filename="d:/tmp/y.ff")
z <- ff(sample(255, n, TRUE), vmode="ubyte", length=n, filename="d:/tmp/z.ff")
df <- ffd(x=x, y=y, z=z)
rm(x,y,z)

message("save all of them")
```

```

ffsave.image("d:/tmp/x")
str(ffinfo("d:/tmp/x"))

message("save some of them with shorter relative pathnames ...")
ffsave(a, b, file="d:/tmp/y", rootpath="d:/tmp")
str(ffinfo("d:/tmp/y"))

message("... and add others later")
ffsave(df, add=TRUE, file="d:/tmp/y", rootpath="d:/tmp")
str(ffinfo("d:/tmp/y"))

message("... and add others later")
system.time(ffsave(a, file="d:/tmp/z", move=TRUE))
ffinfo("d:/tmp/z")

message("let's delete/close/remove all objects")
close(a) # no file anymore, since we moved a into the ffarchive
delete(b, df)
rm(df, a, b, n)
message("prove it")
ls()

message("restore all but ff files in a different directory")
system.time(ffload("d:/tmp/x", rootpath="d:/tmp2"))
lapply(ls(), function(i)filename(get(i)))

delete(a, b, df)
rm(df, a, b)

ffdrop(c("d:/tmp/x", "d:/tmp/y", "d:/tmp/z"))

## End(Not run)

```

ffsort

*Sorting of ff vectors***Description**

Sorting: sort an ff vector – optionally in-place

**Usage**

```

ffsort(x
, aux = NULL
, has.na = TRUE
, na.last = TRUE
, decreasing = FALSE
, inplace = FALSE
, decorate = FALSE
, BATCHBYTES = getOption("ffmaxbytes")

```

```
, VERBOSE = FALSE
)
```

### Arguments

<code>x</code>	an ff vector
<code>aux</code>	NULL or an ff vector of the same type for temporary storage
<code>has.na</code>	boolean scalar telling ffsort whether the vector might contain NAs. <i>Note</i> that you risk a crash if there are unexpected NAs with <code>has.na=FALSE</code>
<code>na.last</code>	boolean scalar telling ffsort whether to sort NAs last or first. <i>Note</i> that 'boolean' means that there is no third option NA as in <a href="#">sort</a>
<code>decreasing</code>	boolean scalar telling ffsort whether to sort increasing or decreasing
<code>inplace</code>	boolean scalar telling ffsort whether to sort the original ff vector (TRUE) or to create a sorted copy (FALSE, the default)
<code>decorate</code>	boolean scalar telling ffsort whether to decorate the returned ff vector with <a href="#">is.sorted</a> and <a href="#">na.count</a> attributes.
<code>BATCHBYTES</code>	maximum number of RAM bytes ffsort should try not to exceed
<code>VERBOSE</code>	cat some info about the sorting

### Details

ffsort tries to sort the vector in-RAM respecting the BATCHBYTES limit. If a fast sort is not possible, it uses a slower in-place sort (shellsort). If in-RAM is not possible, it uses (a yet simple) out-of-memory algorithm. Like [ramsort](#) the in-RAM sorting method is chosen depending on context information. If a key-index sort can be used, ffsort completely avoids merging disk based subsorts. If argument `decorate=TRUE` is used, then `na.count(x)` will return the number of NAs and `is.sorted(x)` will return TRUE if the sort was done with `na.last=TRUE` and `decreasing=FALSE`.

### Value

An ff vector – optionally decorated with [is.sorted](#) and [na.count](#), see argument 'decorate'

### Note

the ff vector may not have a names attribute

### Author(s)

Jens Oehlschlägel

### See Also

[ramsort](#), [fforder](#), [ffdfsort](#)



**Examples**

```

n <- 1e6
x <- ff(c(NA, 999999:1), vmode="double", length=n)
x <- ffsort(x)
x
is.sorted(x)
na.count(x)
x <- ffsort(x, decorate=TRUE)
is.sorted(x)
na.count(x)
x <- ffsort(x, BATCHBYTES=n, VERBOSE=TRUE)

```

---

ffsuitable	<i>Test ff object for suitability</i>
------------	---------------------------------------

---

**Description**

ffsuitable tests whether FF\_RETURN is an `ff` object like FF\_PROTO and having attributes FF\_ATTR.

**Usage**

```

ffsuitable(FF_RETURN, FF_PROTO = NULL, FF_ATTR = list()
, strict.dimorder = TRUE, fail = "warning")
ffsuitable_attris(x)

```

**Arguments**

<code>x</code>	an object from which to extract attributes for comparison
<code>FF_RETURN</code>	the object to be tested for suitability
<code>FF_PROTO</code>	the prototype object which FF_RETURN should match
<code>FF_ATTR</code>	a list of additional attributes dominating those from FF_PROTO
<code>strict.dimorder</code>	if TRUE ffsuitability requires that the dimorders are standard (ascending)
<code>fail</code>	name of a function to be called if not ffsuitable (default <code>warning</code> )

**Value**

TRUE if FF\_RETURN object is suitable, FALSE otherwise

**Note**

xx Work in progress for package **R.ff**

**Author(s)**

Jens Oehlschlägel

**See Also**

[ffconform](#), [ffreturn](#)

---

ffxtensions

*Test for availability of ff extensions*

---

**Description**

checks if this version of package ff supports ff extensions.

**Usage**

```
ffxtensions()  
ffsymmxtensions()
```

**Details**

ff extensions are needed for certain bitcompressed vmodes and ff symm extensions for symmetric matrices.

**Value**

logical scalar

**Author(s)**

Jens Oehlschlägel

**See Also**

[vmode](#)

**Examples**

```
ffxtensions()  
ffsymmxtensions()
```

---

file.resize	<i>Change size of move an existing file</i>
-------------	---

---

### Description

Change size of an existing file (on some platforms sparse files are used) or move file to other name and/or location.

### Usage

```
file.resize(path, size)
file.move(from, to)
```

### Arguments

path	file path (on windows it uses a 'windows' backslash path!)
size	new filesize in bytes as double
from	old file path
to	new file path

### Details

file.resize can enlarge or shrink the file. When enlarged, the file is filled up with zeros. Some platform implementations feature sparse files, so that this operation is very fast. We have tested:

- Ubuntu Linux 8, i386
- FreeBSD 7, i386
- Gentoo Linux Virtual-Server, i386
- Gentoo Linux, x86\_64
- Windows XP

The following work but do not support sparse files

- Mac OS X 10.5, i386
- Mac OS X 10.4, PPC

file.move tries to [file.rename](#), if this fails (e.g. across file systems) the file is copied to the new location and the old file is removed, see [file.copy](#) and [file.remove](#).

### Value

logical scalar representing the success of this operation

### Author(s)

Daniel Adler

**See Also**

[file.create](#), [file.rename](#), [file.info](#), [file.copy](#), [file.remove](#)

**Examples**

```
x <- tempfile()
newsize <- 23 # resize and size to 23 bytes.
file.resize(x, newsize)
file.info(x)$size == newsize
## Not run:
newsize <- 8*(2^30) # create new file and size to 8 GB.
file.resize(x, newsize)
file.info(x)$size == newsize

## End(Not run)
y <- tempfile()
file.move(x,y)
file.remove(y)
```

---

filename

*Get or set filename*

---

**Description**

Get or set filename from ram or `ff` object via the `filename` and `filename<-` generics or rename all files behind a `ffdf` using the `pattern<-` generic.

**Usage**

```
filename(x, ...)
filename(x, ...) <- value
## Default S3 method:
filename(x, ...)
## S3 method for class 'ff_pointer'
filename(x, ...)
## S3 method for class 'ffdf'
filename(x, ...)
## S3 replacement method for class 'ff'
filename(x, ...) <- value
pattern(x, ...)
pattern(x, ...) <- value
## S3 method for class 'ff'
pattern(x, ...)
## S3 replacement method for class 'ff'
pattern(x, ...) <- value
## S3 replacement method for class 'ffdf'
pattern(x, ...) <- value
```

**Arguments**

x	a ram or ff object, or for pattern assignment only - a ffd object
value	a new filename
...	dummy to keep R CMD CHECK quiet

**Details**

Assigning a `filename<-` means renaming the corresponding file on disk - even for ram objects. If that fails, the assignment fails. If a file is moved in or out of `getOption("fftempdir")` the `finalizer` is changed accordingly to 'delete' in `getOption("fftempdir")` and 'close' otherwise. A pattern is an incomplete filename (optional path and optional filename-prefix) that is turned to filenames by adding a random string using and optionally an extension from optionally an extension from `getOption("ffextension")` (see `fftempfile`). `filename<-` exhibits R's standard behaviour of considering "filename" and "./filename" both to be located in `getwd`. By contrast `pattern<-` will create "filename" without path in `getOption("fftempdir")` and only "./filename" in `getwd`.

**Value**

`filename` and `pattern` return a character filename or pattern. For `ffdf` returns a list with one filename element for each `physical` component. The assignment functions return the changed object, which will keep the change even without re-assigning the return-value

**Author(s)**

Jens Oehlschlägel

**See Also**

`fftempfile`, `finalizer`, `ff`, `as.ff`, `as.ram`, `update.ff`, `file.move`

**Examples**

```
## Not run:
message("Neither giving pattern nor filename gives a random filename
with extension ffextension in fftempdir")
x <- ff(1:12)
finalizer(x)
filename(x)
message("Giving a pattern with just a prefix moves to a random filename
beginning with the prefix in fftempdir")
pattern(x) <- "myprefix_"
filename(x)
message("Giving a pattern with a path and prefix moves to a random filename
beginning with prefix in path (use . for getwd) ")
pattern(x) <- "./myprefix"
filename(x)
message("Giving a filename moves to exactly this filename and extension
in the R-expected place) ")
if (!file.exists("./myfilename.myextension")){
```

```

    filename(x) <- "./myfilename.myextension"
    filename(x)
  }

  message("NOTE that the finalizer has changed from 'delete' to 'close':
now WE are responsible for deleting the file - NOT the finalizer")
  finalizer(x)
  delete(x)
  rm(x)

## End(Not run)

```

---

 finalize

*Call finalizer*


---

### Description

This calls the currently assigned finalizer, either via R's finalization mechanism or manually.

### Usage

```

finalize(x, ...)
## S3 method for class 'ff_pointer'
finalize(x, ...)
## S3 method for class 'ff'
finalize(x, ...)
## S3 method for class 'ffdf'
finalize(x, ...)

```

### Arguments

x	either an <a href="#">ff</a> or <a href="#">ffdf</a> object or an <code>ff_pointer</code> , see details
...	currently ignored

### Details

The `finalize.ff_pointer` method is called from R after it had been passed to [reg.finalizer](#). It will set the finalizer name to NULL and call the finalizer.

The `finalize` generic can be called manually on [ff](#) or [ffdf](#) objects. It will call the finalizer but not touch the finalizer name.

For more details see [finalizer](#)

### Value

returns whatever the called finalizer returns, for `ffdf` a list with the finalization returns of each physical component is returned.

**Note**

finalize.ff\_pointer MUST NEVER be called manually - neither directly nor by calling the generic on an ff\_pointer (could erroneously signal that there is no pending finalization lurking around)

**Author(s)**

Jens Oehlschlägel

**See Also**

[finalizer](#)

**Examples**

```
x <- ff(1:12, pattern="./finalizerdemo")
fnam <- filename(x)
finalizer(x)
is.open(x)
file.exists(fnam)

finalize(x)

finalizer(x)
is.open(x)
file.exists(fnam)

delete(x)
finalizer(x)
is.open(x)
file.exists(fnam)

rm(x)
gc()
```

---

finalizer

*Get and set finalizer (name)*

---

**Description**

The generic finalizer allows to get the current finalizer. The generic finalizer<- allows to set the current finalizer or to change an existing finalizer (but not to remove a finalizer).

**Usage**

```
finalizer(x, ...)
finalizer(x, ...) <- value
## S3 method for class 'ff'
finalizer(x, ...)
```

```
## S3 replacement method for class 'ff'
finalizer(x, ...) <- value
```

### Arguments

x	an <code>ff</code> object
value	the name of the new finalizer
...	ignored

### Details

If an `ff` object is created a finalizer is assigned, it has the task to free resources no longer needed, for example remove the `ff` file or free the C++ RAM associated with an open `ff` file. The assigned finalizer depends on the location of the `ff` file: if the file is created in `getOption(fftempdir)` it is considered temporary and has default finalizer `delete`, files created in other locations have default finalizer `close`. The user can override this either by setting `options("fffinalizer")` or by using argument `finalizer` when creating single `ff` objects. Available finalizer generics are "close", "delete" and "deleteIfOpen", available methods are `close.ff`, `delete.ff` and `deleteIfOpen.ff`. In order to be able to change the finalizer before finalization, the finalizer is NOT directly passed to R's finalization mechanism `reg.finalizer` (an active finalizer can never be changed other than be executed). Instead the NAME of the desired finalizer is stored in the `ff` object and `finalize.ff_pointer` is passed to `reg.finalizer`. `finalize.ff_pointer` will at finalization-time determine the desired finalizer and call it.

There are two possible triggers for execution `finalize.ff_pointer`:

1. the garbage collection `gc` following removal `rm` of the `ff` object
2. closing R if `finonexit` was TRUE at `ff` creation-time, determined by `options("fffinonexit")` and `ff` argument `finonexit`

Furthermore there are two possible triggers for calling the finalizer

1. an explicit call to `finalize`
2. an explicit call to one of the finalizers `close`, `delete` and `deleteIfOpen`

The user can define custom finalizers by creating a generic function like `delete`, a `ff_pointer` method like `delete.ff_pointer` and a `ff` method for manual calls like `delete.ff`. The user then is responsible to take care of two things

1. adequate freeing of resources
2. proper maintenance of the finalizer name in the `ff` object via `physical$finalizer`

`is.null(finalizer(ff))` indicates NO active finalizer, i.e. no pending execution of `finalize.ff_pointer` lurking around after call of `reg.finalizer`. This requires that

1. the `ff_pointer` method sets the finalizer name to NULL
2. the `ff` may change a non-NULL finalizer name to a different name but not change it to NULL



**Value**

`finalizer` returns the name of the active finalizer or NULL if no finalizer is active.  
`finalizer<-` returns the changed ff object (reassignment of this return value not needed to keep the change). If there was no pending call to `finalize.ff_pointer` (`is.null(finalizer(ff))`), `finalizer<-` will create one by calling `reg.finalizer` with the current setting of `physical$finonexit`.

**Note**

You can not assign NULL to an active finalizer using `finalizer<-` because this would not stop R's finalization mechanism and would carry the risk of assigning MULTIPLE finalization tasks.

**Author(s)**

Jens Oehlschlägel

**See Also**

`ff`, `finalize`, `reg.finalizer`

**Examples**

```
x <- ff(1:12, pattern="./finalizerdemo")
fnam <- filename(x)
finalizer(x)
finalizer(x) <- "delete"
finalizer(x)
rm(x)
file.exists(fnam)
gc()
file.exists(fnam)
```

---

fixdiag

*Test for fixed diagonal*

---

**Description**

Check if an object has fixed diagonal

**Usage**

```
fixdiag(x, ...)
fixdiag(x, ...) <- value
## S3 method for class 'ff'
fixdiag(x, ...)
## Default S3 method:
fixdiag(x, ...)
## S3 method for class 'dist'
fixdiag(x, ...)
```

**Arguments**

x                    an ff or ram object  
value                assignement value  
...                   further arguments (not used)

**Details**

ff symmetric matrices can be declared to have fixed diagonal at creation time. Compatibility function `fixdiag.default` returns NULL, `fixdiag.dist` returns 0.

**Value**

NULL or the scalar representing the fixed diagonal

**Author(s)**

Jens Oehlschlägel

**See Also**

[fixdiag](#), [ff](#), [dist](#)

**Examples**

```
fixdiag(matrix(1:16, 4, 4))  
fixdiag(dist(rnorm(1:4)))
```

---

geterror.ff

*Get error and error string*

---

**Description**

Get last error code and error string that occurred on an ff object.

**Usage**

```
geterror.ff(x)  
geterrstr.ff(x)
```

**Arguments**

x                    an ff object

**Value**

`geterror.ff` returns an error integer code (no error = 0) and `geterrstr.ff` returns the error message (no error = "no error").

**Author(s)**

Jens Oehlschlägel, Daniel Adler (C++ back-end)

**See Also**

[ff](#)

**Examples**

```
x <- ff(1:12)
geterror.ff(x)
geterrstr.ff(x)
rm(x); gc()
```

---

getpagesize

*Get page size information*

---

**Description**

The function is used for obtaining the natural OS-specific page size in Bytes. `getpagesize` returns the OS-specific page size in Bytes for memory mapped files, while `getdefaultpagesize` returns a suggested page size. `getalignedpagesize` returns the pagesize as a multiple of the OS-specific page size in Bytes, which is the correct way to specify pagesize in `ff`.

**Usage**

```
getpagesize()
getdefaultpagesize()
getalignedpagesize(pagesize)
```

**Arguments**

`pagesize`      a desired pagesize in bytes

**Value**

An integer giving the page size in Bytes.

**Author(s)**

Daniel Adler, Jens Oehlschlägel

**Examples**

```
getpagesize()
getdefaultpagesize()
getalignedpagesize(2000000)
```

---

`getset.ff`*Reading and writing vectors of values (low-level)*

---

### Description

The three functions `get.ff`, `set.ff` and `getset.ff` provide the simplest interface to access an ff file: getting and setting vector of values identified by positive subscripts

### Usage

```
get.ff(x, i)
set.ff(x, i, value, add = FALSE)
getset.ff(x, i, value, add = FALSE)
```

### Arguments

<code>x</code>	an ff object
<code>i</code>	an index position within the ff file
<code>value</code>	the value to write to position <code>i</code>
<code>add</code>	TRUE if the value should rather increment than overwrite at the index position

### Details

`getset.ff` combines the effects of `get.ff` and `set.ff` in a single operation: it retrieves the old value at position `i` before changing it. `getset.ff` will maintain `na.count`.

### Value

`get.ff` returns a vector, `set.ff` returns the 'changed' ff object (like all assignment functions do) and `getset.ff` returns the value at the subscript positions. More precisely `getset.ff(x, i, value, add=FALSE)` returns the old values at the subscript positions `i` while `getset.ff(x, i, value, add=TRUE)` returns the incremented values at the subscript positions.

### Note

`get.ff`, `set.ff` and `getset.ff` are low level functions that do not support `ramclass` and `ramattribs` and thus will not give the expected result with `factor` and `POSIXct`

### Author(s)

Jens Oehlschlägel

### See Also

[readwrite.ff](#) for low-level access to contiguous chunks and [\[.ff](#) for high-level access

**Examples**

```
x <- ff(0, length=12)
get.ff(x, 3L)
set.ff(x, 3L, 1)
x
set.ff(x, 3L, 1, add=TRUE)
x
getset.ff(x, 3L, 1, add=TRUE)
getset.ff(x, 3L, 1)
x
rm(x); gc()
```

---

hi

*Hybrid index class*


---

**Description**

Class for hybrid index representation, plain and rle-packed

**Usage**

```
hi(from, to, by = 1L, maxindex = NA, vw = NULL, pack = TRUE, NAs = NULL)
## S3 method for class 'hi'
print(x, ...)
## S3 method for class 'hi'
str(object, nest.lev=0, ...)
```

**Arguments**

from	integer vector of lower sequence bounds
to	integer vector of upper sequence bounds
by	integer of stepsizes
maxindex	maximum indep position (needed for negative indices)
vw	virtual window information, see <a href="#">vw</a>
pack	FALSE to suppress rle-packing
NAs	a vector of NA positions (not yet used)
x	an object of class 'hi' to be printed
object	an object of class 'hi' to be str'ed
nest.lev	current nesting level in the recursive calls to str
...	further arguments passed to the next method

## Details

Class `hi` will represent index data either as a plain positive or negative index vector or as an rle-packed version thereof. The current implementation switches from plain index positions `i` to rle-packed storage of `diff(i)` as soon as the compression ratio is 3 or higher. Note that sequences shorter than 2 must never be packed (could cause C-side crash). Furthermore hybrid indices are guaranteed to be sorted ascending, which helps `ffs` access method avoiding to swap repeatedly over the same memory pages (or file positions).

## Value

A list of class 'hi' with components

<code>x</code>	directly accessed by the C-code: the sorted index of class 'rlepack' as returned by <a href="#">rlepack</a>
<code>ix</code>	NULL or positions to restore original order
<code>re</code>	logical scalar indicating if sequence was reversed from descending to ascending (in this case <code>is.null(ix)</code> )
<code>minindex</code>	directly accessed by the C-code: represents the lowest positive subscript to be enumerated in case of negative subscripts
<code>maxindex</code>	directly accessed by the C-code: represents the highest positive subscript to be enumerated in case of negative subscripts
<code>length</code>	number of subscripts, whether negative or positive, not the number of selected elements
<code>dim</code>	NULL or <code>dim</code> – used by <a href="#">as.matrix.hi</a>
<code>dimorder</code>	NULL or <code>dimorder</code>
<code>symmetric</code>	logical scalar indicating whether we have a symmetric matrix
<code>fixdiag</code>	logical scalar indicating whether we have a fixed diagonal (can only be true for symmetric matrices)
<code>vw</code>	virtual window information <a href="#">vw</a>
<code>NAs</code>	NULL or NA positions as returned by <a href="#">rlepack</a>

## Note

`hi` defines the class structure, however usually [as.hi](#) is used to actually Hybrid Index Preprocessing for [ff](#)

## Author(s)

Jens Oehlschlägel

## See Also

[as.hi](#) for coercion, [rlepack](#), [intrle](#), [maxindex](#), [poslength](#)

## Examples

```
hi(c(1, 11, 29), c(9, 19, 21), c(1,1,-2))
as.integer(hi(c(1, 11, 29), c(9, 19, 21), c(1,1,-2)))
```

---

hiparse	<i>Hybrid Index, parsing</i>
---------	------------------------------

---

### Description

hiparse implements the parsing done in Hybrid Index Preprocessing in order to avoid RAM for expanding index expressions. *Not to be called directly*

### Usage

```
hiparse(x, envir, first = as.integer(NA), last = as.integer(NA))
```

### Arguments

x	an index expression, precisely: <a href="#">call</a>
envir	the environment in which to evaluate components of the index expression
first	first index position found so far
last	last index position found so far

### Details

This primitive parser recognises the following tokens: numbers like 1, symbols like x, the colon sequence operator `:` and the concat operator `c`. hiparse will [Recall](#) until the index expression is parsed or an unknown token is found. If an unknown token is found, hiparse evaluates it, inspects it and either accepts it or throws an error, caught by [as.hi.call](#), which falls back to evaluating the index expression and dispatching (again) an appropriate [as.hi](#) method. Reasons for suspending the parsing: if the inspected token is of class `'hi'`, `'ri'`, `'bit'`, `'bitwhich'`, `'is.logical'`, `'is.character'`, `'is.matrix'` or has length>16.

### Value

undefined (and redefined as needed by [as.hi.call](#))

### Author(s)

Jens Oehlschlägel

### See Also

[hi](#), [as.hi.call](#)

`is.ff`*Test for class ff*

---

**Description**

checks if x inherits from class "ff"

**Usage**`is.ff(x)`**Arguments**

x                    any object

**Value**

logical scalar

**Author(s)**

Jens Oehlschlägel

**See Also**

[inherits](#), [as.ff](#), [is.ffdf](#)

**Examples**

```
is.ff(integer())
```

---

`is.ffdf`*Test for class ff*

---

**Description**

checks if x inherits from class "ffdf"

**Usage**`is.ffdf(x)`**Arguments**

x                    any object



**Value**

logical scalar

**Author(s)**

Jens Oehlschlägel

**See Also**

[inherits](#), [as.ffdf](#), [is.ff](#)

**Examples**

```
is.ffdf(integer())
```

---

is.open

*Test if object is opened*

---

**Description**

Test whether an ff or ffdof object or a ff\_pointer is opened.

**Usage**

```
is.open(x, ...)
## S3 method for class 'ff'
is.open(x, ...)
## S3 method for class 'ffdf'
is.open(x, ...)
## S3 method for class 'ff_pointer'
is.open(x, ...)
```

**Arguments**

x                    an ff or ffdof object  
 ...                  further arguments (not used)

**Details**

ff objects open automatically if accessed while closed. For ffdof objects we test all of their [physical](#) components including their [row.names](#) if they are [is.ff](#)

**Value**

TRUE or FALSE (or NA if not all components of an ffdof object are opened or closed)

**Author(s)**

Jens Oehlschlägel

**See Also**[is.readonly](#), [open.ff](#), [close.ff](#)**Examples**

```
x <- ff(1:12)
is.open(x)
close(x)
is.open(x)
rm(x); gc()
```

---

`is.readonly`*Get readonly status*

---

**Description**

Get readonly status of an ff object

**Usage**

```
is.readonly(x, ...)
## S3 method for class 'ff'
is.readonly(x, ...)
```

**Arguments**

x	x
...	...

**Details**

ff objects can be created/opened with `readonly=TRUE`. After each opening of the ff file readonly status is stored in the [physical](#) attributes and serves as the default for the next opening. Thus querying a closed ff object gives the last readonly status.

**Value**

logical scalar

**Author(s)**

Jens Oehlschlägel

**See Also**[open.ff](#), [physical](#)**Examples**

```
x <- ff(1:12)
is.readonly(x)
close(x)
open(x, readonly=TRUE)
is.readonly(x)
close(x)
is.readonly(x)
rm(x)
```

is.sorted

*Getting and setting 'is.sorted' physical attribute***Description**

Functions to mark an ff or ram object as 'is.sorted' and query this. Responsibility to maintain this attribute is with the user.

**Usage**

```
## Default S3 method:
is.sorted(x, ...)
## Default S3 replacement method:
is.sorted(x, ...) <- value
```

**Arguments**

x	an ff or ram object
...	ignored
value	NULL (to remove the 'is.sorted' attribute) or TRUE or FALSE

**Details**

Sorting is slow, see [sort](#). Checking whether an object is sorted can avoid unnecessary sorting – see [is.unsorted](#), [intisasc](#) – but still takes too much time with large objects stored on disk. Thus it makes sense to maintain an attribute, that tells us whether sorting can be skipped. Note that – though you change it yourself – `is.sorted` is a [physical](#) attribute of an object, because it represents an attribute of the *data*, which is shared between different [virtual](#) views of the object.

**Value**

TRUE (if set to TRUE) or FALSE (if set to NULL or FALSE)

**Note**

ff will set `is.sorted(x) <- FALSE` if `clone` or `length<- .ff` have increased length.

**Author(s)**

Jens Oehlschlägel

**See Also**

[is.ordered.ff](#) for testing factor levels, [is.unsorted](#) for testing the data, [intisasc](#) for a quick version thereof, [na.count](#) for yet another `physical` attribute

**Examples**

```
x <- 1:12
is.sorted(x) <- !( is.na(is.unsorted(x)) || is.unsorted(x))
is.sorted(x)
x[1] <- 100L
message("don't forget to maintain once it's no longer TRUE")
is.sorted(x) <- FALSE
message("check whether as 'is.sorted' attribute is maintained")
!is.null(physical(x)$is.sorted)
message("remove the 'is.sorted' attribute")
is.sorted(x) <- NULL
message("NOTE that querying 'is.sorted' still returns FALSE")
is.sorted(x)
```

---

length.ff

*Getting and setting length*

---

**Description**

Gets and sets length of ff objects.

**Usage**

```
## S3 method for class 'ff'
length(x)
## S3 replacement method for class 'ff'
length(x) <- value
```

**Arguments**

x	object to query
value	new object length

**Details**

Changing the length of ff objects is only allowed if no `vw` is used. Changing the length of ff objects will remove any `dim.ff` and `dimnames.ff` attribute. Changing the length of ff objects will remove any `na.count` or `is.sorted` attribute and warn about this. New elements are usually zero, but it may depend on OS and filesystem what they really are. If you want standard R behaviour: filling with NA ,you need to do this yourself. As an exception to this rule, ff objects with `names.ff` will be filled with NA's automatically, and the length of the names will be adjusted (filled with position numbers where needed, which can easily consume a lot of RAM, therefore removing 'names' will help to faster increase length without RAM problems).

**Value**

Integer scalar

**Note**

Special care needs to be taken with regard ff objects that represent factors. For ff factors based on UNSIGNED `vmodes`, new values of zero are silently interpreted as the first factor level. For ff factors based on SIGNED `vmodes`, new values of zero result in illegal factor levels. See `nrow<-`.

**Author(s)**

Jens Oehlschlägel

**See Also**

[length](#), [maxlength](#), [file.resize](#), [dim](#), [virtual](#)

**Examples**

```
x <- ff(1:12)
maxlength(x)
length(x)
length(x) <- 10
maxlength(x)
length(x)
length(x) <- 16
maxlength(x)
length(x)
rm(x); gc()
```

---

length.ffdf

*Getting length of a ffdf dataframe*

---

**Description**

Getting "length" (number of columns) of a ffdf dataframe

**Usage**

```
## S3 method for class 'ffdf'
length(x)
```

**Arguments**

x                    an `ffdf` object

**Value**

integer number of columns

**Author(s)**

Jens Oehlschlägel

**See Also**

[dim.ffdf](#), [length.ff](#), [ffdf](#)

**Examples**

```
length(as.ffdf(data.frame(a=1:26, b=letters)))
gc()
```

---

length.hi

*Hybrid Index, querying*

---

**Description**

Functions to query some index attributes

**Usage**

```
## S3 method for class 'hi'
length(x)
maxindex(x, ...)
poslength(x, ...)
## S3 method for class 'hi'
maxindex(x, ...)
## S3 method for class 'ri'
maxindex(x, ...)
## S3 method for class 'bit'
maxindex(x, ...)
## S3 method for class 'bitwhich'
maxindex(x, ...)
## S3 method for class 'logical'
maxindex(x, ...)
```

```
## Default S3 method:
maxindex(x, ...)
## S3 method for class 'hi'
poslength(x, ...)
## S3 method for class 'ri'
poslength(x, ...)
## S3 method for class 'bit'
poslength(x, ...)
## S3 method for class 'bitwhich'
poslength(x, ...)
## S3 method for class 'logical'
poslength(x, ...)
## Default S3 method:
poslength(x, ...)
```

### Arguments

x	an object of class <a href="#">hi</a>
...	further arguments (not used)

### Details

length.hi returns the number of the subscript elements in the index (even if they are negative). By contrast the generic poslength returns the number of selected elements (which for negative indices is `maxindex(x) - length(unique(x))`). The generic maxindex returns the highest possible index position.

### Value

an integer scalar

### Note

duplicated negative indices are removed

### Author(s)

Jens Oehlschlägel

### See Also

[hi](#), [as.hi](#), [length.ff](#), [length](#)

### Examples

```
length(as.hi(-1, maxindex=12))
poslength(as.hi(-1, maxindex=12))
maxindex(as.hi(-1, maxindex=12))
message("note that")
length(as.hi(c(-1, -1), maxindex=12))
length(as.hi(c(1,1), maxindex=12))
```

---

 levels.ff

*Getting and setting factor levels*


---

### Description

levels.ff<- sets factor levels, levels.ff gets factor levels

### Usage

```
## S3 method for class 'ff'
levels(x)
## S3 replacement method for class 'ff'
levels(x) <- value
  is.factor(x)
  is.ordered(x)
## S3 method for class 'ff'
is.factor(x)
## S3 method for class 'ff'
is.ordered(x)
## Default S3 method:
is.factor(x)
## Default S3 method:
is.ordered(x)
```

### Arguments

x	an ff object
value	the new factor levels, if NA is an allowed level it needs to be given explicitly, nothing is excluded

### Details

The ff object must have an integer vmode, see [.rammode](#). If the mode is unsigned – see [.vunsigned](#) – the first factor level is coded with 0L instead of 1L in order to maximize the number of codable levels. Usually the internal ff coding – see [ram2ffcode](#) – is invisible to the user: when subscripting from an ff factor, unsigned codings are automatically converted to R’s standard factor codes starting at 1L. However, you need to be aware of the internal ff coding in two situations.

1. If you convert an ff integer object to an ff factor object and vice versa by assigning levels and `is.null(oldlevels)!=is.null(newlevels)`.
2. Assigning data that does not match any level usually results in NA, however, in unsigned types there is no NA and all unknown data are mapped to the first level.

### Value

levels returns a character vector of levels (possibly including `as.character(NA)`).



**Note**

When levels are assigned to an ff object that formerly had not levels, we assign automatically `ramclass == "factor"`. If you want to change to an ordered factor, use `virtual$ramclass <- c("ordered", "factor")`

**Author(s)**

Jens Oehlschlägel

**See Also**

[ramclass](#), [factor](#), [virtual](#)

**Examples**

```
message("--- create an ff factor including NA as last level")
x <- ff("a", levels=c(letters, NA), length=99)
message(' we expect a warning because "A" is an unknown level')
x[] <- c("a", NA,"A")
x
levels(x)

message("--- create an ff ordered factor")
x <- ff(letters, levels=letters, ramclass=c("ordered","factor"), length=260)
x
levels(x)

message(" make it a non-ordered factor")
virtual(x)$ramclass <- "factor"
x
rm(x); gc()

## Not run:
message("--- create an unsigned quad factor")
x <- ff(c("A","T","G","C"), levels=c("A","T","G","C"), vmode="quad", length=100)
x
message(" 0:3 coding usually invisible to the user")
unclass(x[1:4])
message(" after removing levels, the 0:3 coding becomes visible to the user")
message(" we expect a warning here")
levels(x) <- NULL
x[1:4]
rm(x); gc()

## End(Not run)
```

## Description

This help page lists the currently known limitations of package `ff`, as well as differences between `ff` and `ram` methods.

### Automatic file removal

Remind that not giving parameter `ff(filename=)` will result in a temporary file in `fftempdir` with 'delete' finalizer, while giving parameter `ff(filename=)` will result in a permanent file with 'close' finalizer. Do avoid setting `setwd(getOption("fftempdir"))!` Make sure you really understand the implications of automatic unlinking of `getOption("fftempdir")` `.onUnload`, of finalizer choice and of finalizing behaviour at the end of R sessions as defaulted in `getOption("ffinonexit")`. **Otherwise you might experience 'unexpected' losses of files and data.**

### Size of objects

Currently `ff` objects cannot have length zero and are limited to `.Machine$integer.max` elements. We have not yet ported the R code to support 64bit double indices (in essence 52 bits integer) although the C++ back-end has been prepared for this. Furthermore filesize limitations of the OS apply, see `ff`.

### Side effects

In contrast to standard R expressions, `ff` expressions violate the functional programming logic and are called for their side effects. This is also true for `ram` compatibility functions `swap.default`, and `add.default`.

### Hybrid copying semantics

If you modify a copy of an `ff` object, changes of data (`[<-`) and of `physical` attributes will be shared, but changes in `virtual` and class attributes will not.

### Limits of compatibility between `ff` and `ram` objects

If it's not too big, you can move an `ff` object completely into R's RAM through `as.ram`. However, you should watch out for three limitations:

1. `Ram` objects don't have hybrid copying semantics; changes to a copy of a `ram` object will never change the original `ram` object
2. Assigning values to a `ram` object can easily upgrade to a higher `storage.mode`. This will create conflicts with the `vmode` of the `ram` object, which goes undetected until you try to write back to disk through `as.ff`.
3. Writing back to disk with `as.ff` under the same filename requires that the original `ff` object has been deleted (or at least closed if you specify parameter `overwrite=TRUE`).

**Index expressions**

ff index expressions do not allow zeros and NAs, see [\[.ff](#) and see [as.hi](#)

**Availability of bydim parameter**

Parameter bydim is only available in ff access methods, see [\[.ff](#)

**Availability of add parameter**

Parameter add is only available in ff access methods, see [\[.ff](#)

**Compatibility of swap and add**

If index expressions contain duplicated positions, the ff and ram methods for [swap](#) and [add](#) will behave differently, see [swap](#).

**Definition of [[ and [[<-**

You should consider the behaviour of [\[\[.ff](#) and [\[\[<-.ff](#) as undefined and not use them in programming. Currently they are shortcuts to [get.ff](#) and [set.ff](#), which unlike [\[.ff](#) and [\[<-.ff](#) do not support [factor](#) and [POSIXct](#), nor [dimorder](#) or virtual windows [vw](#). In contrast to the standard methods, [\[\[.ff](#) and [\[\[<-.ff](#) only accepts positive integer index positions. The definition of [\[\[.ff](#) and [\[\[<-.ff](#) may be changed in the future.

**Multiple vector interpretation in arrays**

R objects have always standard [dimorder](#) `1:length(dim)`. In case of non-standard [dimorder](#) (see [dimorderStandard](#)) the vector sequence of array elements in R and in the ff file differs. To access array elements in file order, you can use [getset.ff](#), [readwrite.ff](#) or copy the ff object and set `dim(ff)<-NULL` to get a vector view into the ff object (using `[` dispatches the vector method [\[.ff](#)). To access the array elements in R standard [dimorder](#) you simply use `[` which dispatches to [\[.ff\\_array](#). Note that in this case [as.hi](#) will unpack the complete index, see next section.

**RAM expansion of index expressions**

Some index expressions do not consume RAM due to the [hi](#) representation. For example `1:n` will almost consume no RAM however large `n`. However, some index expressions are expanded and require to `maxindex(i) * .rambytes["integer"]` bytes, either because the sorted sequence of index positions cannot be rle-packed efficiently or because [hiparse](#) cannot yet parse such expression and falls back to evaluating/expanding the index expression. If the index positions are not sorted, the index will be expanded and a second vector is needed to store the information for re-ordering, thus the index requires `2 * maxindex(i) * .rambytes["integer"]` bytes.

**RAM expansion when recycling assignment values**

Some assignment expressions do not consume RAM for recycling. For example `x[1:n] <- 1:k` will not consume RAM however large is `n` compared to `k`, when `x` has standard [dimorder](#). However, if `length(value)>1`, assignment expressions with non-ascending index positions trigger recycling the value R-side to the full index length. This will happen if [dimorder](#) does not match parameter [bydim](#) or if the index is not sorted in ascending order.

**Byteorder incompatibility**

Note that ff files cannot be transferred between systems with different byteorder.

---

matcomb

*Array: make matrix indices from row and columns positions*

---

**Description**

create matrix indices from row and columns positions

**Usage**

```
matcomb(r, c)
```

**Arguments**

r                    integer vector of row positions  
c                    integer vector of column positions

**Details**

rows rotate faster than columns

**Value**

a k by 2 matrix of matrix indices where  $k = \text{length}(r) * \text{length}(c)$

**Author(s)**

Jens Oehlschlägel

**See Also**

[row](#), [col](#), [expand.grid](#)

**Examples**

```
matcomb(1:3, 1:4)  
matcomb(2:3, 2:4)
```

---

matprint	<i>Print beginning and end of big matrix</i>
----------	--

---

## Description

Print beginning and end of big matrix

## Usage

```
matprint(x, maxdim = c(16, 16), digits = getOption("digits"))  
## S3 method for class 'matprint'  
print(x, quote = FALSE, right = TRUE, ...)
```

## Arguments

x	a <a href="#">matrix</a>
maxdim	max number of rows and columns for printing
digits	see <a href="#">format</a>
quote	see <a href="#">print</a>
right	see <a href="#">print</a>
...	see <a href="#">print</a>

## Value

a list of class 'matprint' with components

subscript	a list with four vectors of subscripts: row begin, column begin, row end, column end
example	the extracted example matrix as character including separators
rsep	logical scalar indicating whether row separator is included
csep	logical scalar indicating whether column separator is included

## Author(s)

Jens Oehlschlägel

## See Also

[vecprint](#)

## Examples

```
matprint(matrix(1:(300*400), 300, 400))
```

---

maxffmode	<i>Lossless vmode coercability</i>
-----------	------------------------------------

---

**Description**

maxffmode returns the lowest [vmode](#) that can absorb all input vmodes without data loss

**Usage**

```
maxffmode(...)
```

**Arguments**

...                   one or more vectors of vmodes

**Value**

the smallest [.ffmode](#) which can absorb the input vmodes without data loss

**Note**

The output can be larger than any of the inputs (if the highest input vmode is an integer type without NA and any other input requires NA).

**Author(s)**

Jens Oehlschlägel

**See Also**

[.vcoerceable](#), [.ffmode](#), [ffconform](#)

**Examples**

```
maxffmode(c("quad", "logical"), "ushort")
```

---

maxlength	<i>Get physical length of an ff or ram object</i>
-----------	---

---

**Description**

maxlength returns the physical length of an ff or ram object

**Usage**

```
maxlength(x, ...)  
## S3 method for class 'ff'  
maxlength(x, ...)  
## Default S3 method:  
maxlength(x, ...)
```

**Arguments**

x	ff or ram object
...	additional arguments (not used)

**Value**

integer scalar

**Author(s)**

Jens Oehlschlägel

**See Also**

[length.ff](#), [maxindex](#)

**Examples**

```
x <- ff(1:12)  
length(x) <- 10  
length(x)  
maxlength(x)  
x  
rm(x); gc()
```

---

`mismatch`*Test for recycle mismatch*

---

**Description**

`mismatch` will return TRUE if the larger of `nx,ny` is not a multiple of the other and the other is  $>0$  (see `arithmetic.c`). `ymismatch` will return TRUE if `nx` is not a multiple of `ny` and `ny`  $>0$

**Usage**

```
mismatch(nx, ny)
ymismatch(nx, ny)
```

**Arguments**

<code>nx</code>	x length
<code>ny</code>	y length

**Value**

logical scalar

**Author(s)**

Jens Oehlschlägel

**See Also**

[ffconform](#)

**Examples**

```
ymismatch(4,0)
ymismatch(4,2)
ymismatch(4,3)
ymismatch(2,4)
mismatch(4,0)
mismatch(4,2)
mismatch(4,3)
mismatch(2,4)
```



---

`na.count`*Getting and setting 'na.count' physical attribute*

---

### Description

The 'na.count' physical attribute gives the current number of NAs if properly initialized and properly maintained, see details.

### Usage

```
## S3 method for class 'ff'  
na.count(x, ...)  
## Default S3 method:  
na.count(x, ...)  
## S3 replacement method for class 'ff'  
na.count(x, ...) <- value  
## Default S3 replacement method:  
na.count(x, ...) <- value
```

### Arguments

<code>x</code>	an ff or ram object
<code>...</code>	further arguments (not used)
<code>value</code>	NULL (to remove the 'na.count' attribute) or TRUE to activate or an integer value

### Details

The 'na.count' feature is activated by assigning the current number of NAs to `na.count(x) <- currentNA` and deactivated by assigning NULL. The 'na.count' feature is maintained by the, [getset.ff](#), [readwrite.ff](#) and [swap](#), other ff methods for writing – [set.ff](#), [\[\[<-.ff](#), [write.ff](#), [\[<-.ff](#) – will stop if 'na.count' is activated. The functions `na.count` and `na.count<-` are generic. For ram objects, the default method for `na.count` calculates the number of NAs on the fly, thus no maintenance restrictions apply.

### Value

NA (if set to NULL or NA) or an integer value otherwise

### Author(s)

Jens Oehlschlägel, Daniel Adler (C++ back-end)

### See Also

[getset.ff](#), [readwrite.ff](#) and [swap](#) for methods that support maintenance of 'na.count', [NA](#), [is.sorted](#) for yet another [physical](#) attribute

**Examples**

```

message("--- ff examples ---")
x <- ff(1:12)
na.count(x)
message("activate the 'na.count' physical attribute and set the current na.count manually")
na.count(x) <- 0L
message("add one NA with a method that maintains na.count")
swap(x, NA, 1)
na.count(x)
message("remove the 'na.count' physical attribute (and stop automatic maintenance)")
na.count(x) <- NULL
message("activate the 'na.count' physical attribute and have ff automatically
calculate the current na.count")
na.count(x) <- TRUE
na.count(x)
message("--- ram examples ---")
x <- 1:12
na.count(x)
x[1] <- NA
message("activate the 'na.count' physical attribute and have R automatically
calculate the current na.count")
na.count(x) <- TRUE
na.count(x)
message("remove the 'na.count' physical attribute (and stop automatic maintenance)")
na.count(x) <- NULL
na.count(x)
rm(x); gc()

```

---

names.ff

*Getting and setting names*


---

**Description**

For `ff_vectors` you can set names, though this is not recommended for large objects.

**Usage**

```

## S3 method for class 'ff'
names(x)
## S3 replacement method for class 'ff'
names(x) <- value
## S3 method for class 'ff_array'
names(x)
## S3 replacement method for class 'ff_array'
names(x) <- value

```

**Arguments**

<code>x</code>	a <code>ff</code> vector
<code>value</code>	a character vector

### Details

If `vw` is set, `names.ff` returns the appropriate part of the names, but you can't set names while `vw` is set. `names.ff\_array` returns `NULL` and setting names for `ff\_arrays` is not allowed, but setting `dimnames` is.

### Value

`names` returns a character vector (or `NULL`)

### Author(s)

Jens Oehlschlägel

### See Also

[names](#), [dimnames.ff\\_array](#), [vw](#), [virtual](#)

### Examples

```
x <- ff(1:26, names=letters)
names(x)
names(x) <- LETTERS
names(x)
names(x) <- NULL
names(x)
rm(x); gc()
```

---

nrowAssign

*Assigning the number of rows or columns*

---

### Description

Function `nrow<-` assigns `dim` with a new number of rows.  
Function `ncol<-` assigns `dim` with a new number of columns.

### Usage

```
nrow(x) <- value
ncol(x) <- value
```

### Arguments

<code>x</code>	a object that has <code>dim</code> AND can be assigned ONE new dimension
<code>value</code>	the new size of the assigned dimension

### Details

Currently only assigning new rows to `ffdf` is supported. The new `ffdf` rows are not initialized (usually become zero). NOTE that

**Value**

The object with a modified dimension

**Author(s)**

Jens Oehlschlägel

**See Also**

[ffdf](#), [dim.ffdf](#)

**Examples**

```
a <- as.ff(1:26)
b <- as.ff(factor(letters)) # vmode="integer"
c <- as.ff(factor(letters), vmode="ubyte")
df <- ffdof(a,b,c)
nrow(df) <- 2*26
df
message("NOTE that the new rows have silently the first level 'a' for UNSIGNED vmodes")
message("NOTE that the new rows have an illegal factor level <0> for SIGNED vmodes")
message("It is your responsibility to put meaningful content here")
message("As an example we replace the illegal zeros by NA")
df$b[27:52] <- NA
df

rm(a,b,c,df); gc()
```

---

open.ff

*Opening an ff file*

---

**Description**

open.ff opens an ff file, optionally marking it readonly and optionally specifying a caching scheme.

**Usage**

```
## S3 method for class 'ff'
open(con, readonly = FALSE, pagesize = NULL, caching = NULL, assert = FALSE, ...)
## S3 method for class 'ffdf'
open(con, readonly = FALSE, pagesize = NULL, caching = NULL, assert = FALSE, ...)
```

**Arguments**

con	an <a href="#">ff</a> or <a href="#">ffdf</a> object
readonly	readonly
pagesize	number of bytes to use as pagesize or NULL to take the pagesize stored in the <a href="#">physical</a> attribute of the ff object, see <a href="#">getalignedpagesize</a>

caching            one of 'mnmoflush' or 'mmeachflush', see [ff](#)  
 assert            setting this to TRUE will give a message if the ff was not open already  
 ...                further arguments (not used)

### Details

ff objects will be opened automatically when accessing their content and the file is still closed. Opening `ffdf` objects will open all of their [physical](#) components including their `row.names` if they are `is.ff`

### Value

TRUE if object could be opened, FALSE if it was opened already (or NA if not all components of an `ffdf` returned FALSE or TRUE on opening)

### Author(s)

Jens Oehlschlägel

### See Also

[ff](#), [close.ff](#), [delete](#), [deleteIfOpen](#), [getalignedpagesize](#)

### Examples

```

x <- ff(1:12)
close(x)
is.open(x)
open(x)
is.open(x)
close(x)
is.open(x)
x[]
is.open(x)
y <- x
close(y)
is.open(x)
rm(x,y); gc()

```

---

pagesize

*Pagesize of ff object*

---

### Description

Returns current pagesize of ff object

**Usage**

```
pagesize(x, ...)  
## S3 method for class 'ff'  
pagesize(x, ...)
```

**Arguments**

```
x          an ff object  
...       further arguments (not used)
```

**Value**

integer number of bytes

**Author(s)**

Jens Oehlschlägel

**See Also**

[getpagesize](#)

**Examples**

```
x <- ff(1:12)  
pagesize(x)
```

---

physical.ff

*Getting and setting physical and virtual attributes of ff objects*

---

**Description**

Functions for getting and setting physical and virtual attributes of ff objects.

**Usage**

```
## S3 method for class 'ff'  
physical(x)  
## S3 method for class 'ff'  
virtual(x)  
## S3 replacement method for class 'ff'  
physical(x) <- value  
## S3 replacement method for class 'ff'  
virtual(x) <- value
```

**Arguments**

x                    an ff object  
value                a list with named elements

**Details**

ff objects have physical and virtual attributes, which have different copying semantics: physical attributes are shared between copies of ff objects while virtual attributes might differ between copies. [as.ram](#) will retain some physical and virtual attributes in the ram clone, such that [as.ff](#) can restore an ff object with the same attributes.

**Value**

physical and virtual returns a list with named elements

**Author(s)**

Jens Oehlschlägel

**See Also**

[physical](#), [physical.ffdf](#), [ff](#), [as.ram](#);  
[is.sorted](#) and [na.count](#) for applications of physical attributes;  
[levels.ff](#) and [ramattribs](#) for applications of virtual attributes

**Examples**

```
x <- ff(1:12)
x
physical(x)
virtual(x)
y <- as.ram(x)
physical(y)
virtual(y)
rm(x,y); gc()
```

---

physical.ffdf

*Getting physical and virtual attributes of ffdf objects*

---

**Description**

Functions for getting physical and virtual attributes of ffdf objects.

**Usage**

```
## S3 method for class 'ffdf'
physical(x)
## S3 method for class 'ffdf'
virtual(x)
```

**Arguments**

x                    an `ffdf` object

**Details**

`ffdf` objects enjoy a complete decoupling of virtual behaviour from physical storage. The physical component is simply a (potentially named) list where each element represents an atomic ff vector or matrix. The virtual component is itself a dataframe, each row of which defines a column of the `ffdf` through a mapping to the physical component.

**Value**

'physical.ffdf' returns a `list` with atomic ff objects.

'virtual.ffdf' returns a `data.frame` with the following columns

VirtualVmode	the <code>vmode</code> of this row (=ffdf column)
ASIs	logical defining the <code>ASIs</code> status of this row (=ffdf column)
VirtualIsMatrix	logical defining whether this row (=ffdf column) represents a matrix
PhysicalIsMatrix	logical reporting whether the corresponding physical element is a matrix
PhysicalElementNo	integer identifying the corresponding physical element
PhysicalFirstCol	integer identifying the first column of the corresponding physical element (1 if it is not a matrix)
PhysicalLastCol	integer identifying the last column of the corresponding physical element (1 if it is not a matrix)

**Author(s)**

Jens Oehlschlägel

**See Also**

`ffdf`, `physical`, `virtual`, `vmode`

**Examples**

```
x <- 1:2
y <- matrix(1:4, 2, 2)
z <- matrix(1:4, 2, 2)

message("Here the y matrix is first converted to single columns by data.frame,
then those columns become ff")
d <- as.ffdf(data.frame(x=x, y=y, z=I(z)))
physical(d)
```



```

virtual(d)

message("Here the y matrix is first converted to ff, and then stored still as matrix
in the ffdof object (although virtually treated as columns of ffdof)")
d <- ffdof(x=as.ff(x), y=as.ff(y), z=I(as.ff(z)))
physical(d)
virtual(d)

message("Apply the usual methods extracting physical attributes")
lapply(physical(d), filename)
lapply(physical(d), vmode)
message("And don't confuse with virtual vmode")
vmode(d)

rm(d); gc()

```

---

print.ff

---

*Print and str methods*


---

## Description

printing ff objects and compactly showing their structure

## Usage

```

## S3 method for class 'ff'
print(x, ...)
## S3 method for class 'ff_vector'
print(x, maxlength = 16, ...)
## S3 method for class 'ff_matrix'
print(x, maxdim = c(16, 16), ...)
## S3 method for class 'ff'
str(object, nest.lev=0, ...)
## S3 method for class 'ffdf'
str(object, nest.lev=0, ...)

```

## Arguments

x	a ff object
object	a ff object
nest.lev	current nesting level in the recursive calls to str
maxlength	max number of elements to print from an ff_vector
maxdim	max number of elements to print from each dimension from an ff_array
...	further arguments to print

## Details

The print methods just print a few exemplary elements from the beginning and end of the dimensions.

**Value**

invisible()

**Author(s)**

Jens Oehlschlägel

**See Also**

[ff](#), [print](#), [str](#)

**Examples**

```
x <- ff(1:10000)
x
print(x, maxlength=30)
dim(x) <- c(100,100)
x
rm(x); gc()
```

---

ram2ffcode

*Factor codings*


---

**Description**

Function `ram2ffcode` creates the *internal* factor codes used by `ff` to store factor levels. Function `ram2ramcode` is a compatibility function used instead if `RETURN_FF==FALSE`.

**Usage**

```
ram2ffcode(value, levels, vmode)
ram2ramcode(value, levels)
```

**Arguments**

value	factor or character vector of values
levels	character vector of factor levels
vmode	one of the integer vmodes in <a href="#">.rammode</a>

**Details**

Factors stored in unsigned vmodes [.vunsigned](#) have their first level represented as 0L instead of 1L.

**Value**

A vector of integer values representing the corresponding factor levels.

**Author(s)**

Jens Oehlschlägel

**See Also**

[factor](#), [levels.ff](#), [vmode](#)

**Examples**

```
ram2ffcode(letters, letters, vmode="byte")
ram2ffcode(letters, letters, vmode="ubyte")
ram2ffcode(letters, letters, vmode="nibble")
message('note that ram2ffcode() does NOT warn that vmode="nibble" cannot store 26 levels')
```

---

ramattribs

*Get ramclass and ramattribs*

---

**Description**

Functions `ramclass` and `ramattribs` return the respective virtual attributes, that determine which class (and attributes) an `ff` object receives when subscripted (or coerced) to `ram`.

**Usage**

```
ramclass(x, ...)
## S3 method for class 'ff'
ramclass(x, ...)
## Default S3 method:
ramclass(x, ...)
ramattribs(x, ...)
## S3 method for class 'ff'
ramattribs(x, ...)
## Default S3 method:
ramattribs(x, ...)
```

**Arguments**

<code>x</code>	<code>x</code>
<code>...</code>	further arguments (not used)

**Details**

`ramclass` and `ramattribs` provide a general mechanism to store atomic classes in `ff` objects, for example [factor](#) – see [levels.ff](#) – and [POSIXct](#), see the example.

**Value**

ramclass returns a character vector with classnames and ramattribs returns a list with names elements just like `attributes`. The vectors `ramclass_excludes` and `ramattribs_excludes` name those attributes, which are not exported from ff to ram objects when using `as.ram`.

**Author(s)**

Jens Oehlschlägel

**See Also**

`ff`, `virtual`, `as.ram`, `levels.ff`, `attributes`, `DateTimeClasses`

**Examples**

```
x <- ff(as.POSIXct(as.POSIXlt(Sys.time(), "GMT")), length=12)
x
ramclass(x)
ramattribs(x)
class(x[])
attributes(x[])
virtual(x)$ramattribs$tzzone = NULL
attributes(x[])
rm(x); gc()
```

---

ramorder.default

*Sorting: order R vector in-RAM and in-place*

---

**Description**

Function `ramorder` will order the input vector in-place (without making a copy) and return the number of NAs found

**Usage**

```
## Default S3 method:
ramorder(x, i, has.na = TRUE, na.last = TRUE, decreasing = FALSE
, stable = TRUE, optimize = c("time", "memory"), VERBOSE = FALSE, ...)
## Default S3 method:
mergeorder(x, i, has.na = TRUE, na.last = TRUE, decreasing = FALSE, ...)
## Default S3 method:
radixorder(x, i, has.na = TRUE, na.last = TRUE, decreasing = FALSE, ...)
## Default S3 method:
keyorder(x, i, keyrange=range(x, na.rm=has.na), has.na = TRUE, na.last = TRUE
, decreasing = FALSE, ...)
## Default S3 method:
shellorder(x, i, has.na = TRUE, na.last = TRUE, decreasing = FALSE, stabilize=FALSE, ...)
```

**Arguments**

x	an atomic R vector
i	a integer vector with a permutation of positions in x (you risk a crash if you violate this)
keyrange	an integer vector with two values giving the smallest and largest possible value in x, note that you should give this explicitly for best performance, relying on the default needs one pass over the data to determine the range
has.na	boolean scalar telling ramorder whether the vector might contain NAs. <i>Note</i> that you risk a crash if there are unexpected NAs with has.na=FALSE
na.last	boolean scalar telling ramorder whether to order NAs last or first. <i>Note</i> that 'boolean' means that there is no third option NA as in <a href="#">order</a>
decreasing	boolean scalar telling ramorder whether to order increasing or decreasing
stable	set to false if stable ordering is not needed (may enlarge the set of ordering methods considered)
optimize	by default ramorder optimizes for 'time' which requires more RAM, set to 'memory' to minimize RAM requirements and sacrifice speed
VERBOSE	cat some info about chosen method
stabilize	Set to TRUE for stabilizing the result of shellorder (for equal keys the order values will be sorted, this only works if i=1:n) to minimize RAM requirements and sacrifice speed
...	ignored

**Details**

Function ramorder is a front-end to a couple of single-threaded ordering algorithms that have been carefully implemented to be fast with and without NAs.

The default is a mergeorder algorithm without copying (Sedgewick 8.4) for integer and double data which requires 2x the RAM of its input vector (character or complex data are not supported). Mergeorder is fast, stable with a reliable runtime.

For integer data longer than a certain length we improve on mergeorder by using a faster LSD radixorder algorithm (Sedgewick 10.5) that uses 2x the RAM of its input vector plus 65536+1 integers.

For booleans, logicals, integers at or below the resolution of smallint and for factors below a certain number of levels we use a key-index order instead of mergeorder or radix order (note that R has a (slower) key-index order in [sort.list](#) available with confusingly named method='radix' but the standard [order](#) does not leverage it for factors (2-11.1). If you call keyorder directly, you should provide a known 'keyrange' directly to obtain the full speed.

Finally the user can request a order method that minimizes memory use at the price of longer computation time with optimize='memory' – currently a shellorder.

**Value**

integer scalar with the number of NAs. This is always 0 with has.na=FALSE

**Note**

This function is called for its side-effects and breaks the functional programming paradigm. Use with care.

**Author(s)**

Jens Oehlschlägel

**References**

Robert Sedgewick (1997). Algorithms in C, Third edition. Addison-Wesley.

**See Also**

[order](#), [fforder](#), [dforder](#), [ramsort](#)

**Examples**

```
n <- 50
x <- sample(c(NA, NA, 1:26), n, TRUE)
order(x)
i <- 1:n
ramorder(x, i)
i
x[i]

## Not run:
message("Note how the datatype influences sorting speed")
n <- 1e7
x <- sample(1:26, n, TRUE)

y <- as.double(x)
i <- 1:n
system.time(ramorder(y, i))

y <- as.integer(x)
i <- 1:n
system.time(ramorder(y, i))

y <- as.short(x)
i <- 1:n
system.time(ramorder(y, i))

y <- factor(letters)[x]
i <- 1:n
system.time(ramorder(y, i))

## End(Not run)
```

---

ramsort.default	<i>Sorting: Sort R vector in-RAM and in-place</i>
-----------------	---

---

## Description

Function `ramsort` will sort the input vector in-place (without making a copy) and return the number of NAs found

## Usage

```
## Default S3 method:
ramsort(x, has.na = TRUE, na.last = TRUE, decreasing = FALSE
, optimize = c("time", "memory"), VERBOSE = FALSE, ...)
## Default S3 method:
mergesort(x, has.na = TRUE, na.last = TRUE, decreasing = FALSE, ...)
## Default S3 method:
radixsort(x, has.na = TRUE, na.last = TRUE, decreasing = FALSE, ...)
## Default S3 method:
keysort(x, keyrange=range(x, na.rm=has.na), has.na = TRUE
, na.last = TRUE, decreasing = FALSE, ...)
## Default S3 method:
shellsort(x, has.na = TRUE, na.last = TRUE, decreasing = FALSE, ...)
```

## Arguments

<code>x</code>	an atomic R vector
<code>keyrange</code>	an integer vector with two values giving the smallest and largest possible value in <code>x</code> , note that you should give this explicitly for best performance, relying on the default needs one pass over the data to determine the range
<code>has.na</code>	boolean scalar telling <code>ramsort</code> whether the vector might contain NAs. <i>Note</i> that you risk a crash if there are unexpected NAs with <code>has.na=FALSE</code>
<code>na.last</code>	boolean scalar telling <code>ramsort</code> whether to sort NAs last or first. <i>Note</i> that 'boolean' means that there is no third option NA as in <a href="#">sort</a>
<code>decreasing</code>	boolean scalar telling <code>ramsort</code> whether to sort increasing or decreasing
<code>optimize</code>	by default <code>ramsort</code> optimizes for 'time' which requires more RAM, set to 'memory' to minimize RAM requirements and sacrifice speed
<code>VERBOSE</code>	cat some info about chosen method
<code>...</code>	ignored

## Details

Function `ramsort` is a front-end to a couple of single-threaded sorting algorithms that have been carefully implemented to be fast with and without NAs.

The default is a mergesort algorithm without copying (Sedgewick 8.4) for integer and double data

which requires 2x the RAM of its input vector (character or complex data are not supported). Mergesort is fast, stable with a reliable runtime.

For integer data longer than a certain length we improve on mergesort by using a faster LSD radix-sort algorithm (Sedgewick 10.5) that uses 2x the RAM of its input vector plus 65536+1 integers.

For booleans, logicals, integers at or below the resolution of `smallint` and for factors below a certain number of levels we use a key-index sort instead of mergesort or radix sort (note that R has a (slower) key-index sort in `sort.list` available with confusingly named `method='radix'` but the standard `sort` does not leverage it for factors (2-11.1). If you call `keysort` directly, you should provide a known 'keyrange' directly to obtain the full speed.

Finally the user can request a sort method that minimizes memory use at the price of longer computation time with `optimize='memory'` – currently a shellsort.

### Value

integer scalar with the number of NAs. This is always 0 with `has.na=FALSE`

### Note

This function is called for its side-effects and breaks the functional programming paradigm. Use with care.

### Author(s)

Jens Oehlschlägel

### References

Robert Sedgewick (1997). Algorithms in C, Third edition. Addison-Wesley.

### See Also

[sort](#), [ffsort](#), [dfsort](#), [ramorder](#)

### Examples

```
n <- 50
x <- sample(c(NA, NA, 1:26), n, TRUE)
sort(x)
ramsort(x)
x

## Not run:
message("Note how the datatype influences sorting speed")
n <- 5e6
x <- sample(1:26, n, TRUE)

y <- as.double(x)
system.time(ramsort(y))

y <- as.integer(x)
system.time(ramsort(y))
```



```

y <- as.short(x)
system.time(ramsort(y))

y <- as.factor(letters)[x]
system.time(ramsort(y))

## End(Not run)

```

---

read.table.ffdf

*Importing csv files into ff data.frames*


---

## Description

Function `read.table.ffdf` reads separated flat files into `ffdf` objects, very much like (and using) `read.table`. It can also work with any convenience wrappers like `read.csv` and provides its own convenience wrapper (e.g. `read.csv.ffdf`) for R's usual wrappers.

## Usage

```

read.table.ffdf(
  x = NULL
  , file, fileEncoding = ""
  , nrows = -1, first.rows = NULL, next.rows = NULL
  , levels = NULL, appendLevels = TRUE
  , FUN = "read.table", ...
  , transFUN = NULL
  , asffdf_args = list()
  , BATCHBYTES = getOption("ffbatchbytes")
  , VERBOSE = FALSE
)
read.csv.ffdf(...)
read.csv2.ffdf(...)
read.delim.ffdf(...)
read.delim2.ffdf(...)

```

## Arguments

- |      |   |
|------|---|
| x    | NULL or an optional <code>ffdf</code> object to which the read records are appended. If this is provided, it defines crucial features that are otherwise determined during the 'first' chunk of reading: <code>vmodes</code> , <code>colnames</code> , <code>colClasses</code> , sequence of predefined <code>levels</code> . In order to also read the first chunk into such predefined <code>ffdf</code> , an x with 1 row is treated special: instead of appending the first row will be overwritten. This is necessary because we cannot provide x with zero rows (we cannot create ff vectors with zero elements). |
| file | the name of the file which the data are to be read from. Each row of the table appears as one line of the file. If it does not contain an <i>absolute</i> path, the file  |

	name is <i>relative</i> to the current working directory, <code>getwd()</code> . Tilde-expansion is performed where supported.
	Alternatively, <code>file</code> can be a readable text-mode <code>connection</code> (which will be opened for reading if necessary, and if so <code>closed</code> (and hence destroyed) at the end of the function call).
<code>fileEncoding</code>	character string: if non-empty declares the encoding used on a file (not a connection) so the character data can be re-encoded. See <code>file</code> .
<code>nrows</code>	integer: the maximum number of rows to read in (includes <code>first.rows</code> in case a 'first' chunk is read) Negative and other invalid values are ignored.
<code>first.rows</code>	integer: number of rows to be read in the first chunk, see details. Default is the value given at <code>next.rows</code> or <code>1e3</code> otherwise. Ignored if <code>x</code> is given.
<code>next.rows</code>	integer: number of rows to be read in further chunks, see details. By default calculated as <code>BATCHBYTES %% sum(.rambytes[vmode(x)])</code>
<code>levels</code>	NULL or an optional list, each element named with <code>col.names</code> of factor columns specifies the <code>levels</code> Ignored if <code>x</code> is given.
<code>appendLevels</code>	logical. A vector of permissions to expand <code>levels</code> for factor columns. Recycled as necessary, or if the logical vector is named, unspecified values are taken to be TRUE. Ignored during processing of the 'first' chunk
<code>FUN</code>	character: name of a function that is called for reading each chunk, see <code>read.table</code> , <code>read.csv</code> , etc.
<code>...</code>	further arguments, passed to <code>FUN</code> in <code>read.table.ffdf</code> , or passed to <code>read.table.ffdf</code> in the convenience wrappers
<code>transFUN</code>	NULL or a function that is called on each data.frame chunk after reading with <code>FUN</code> and before further processing (for filtering, transformations etc.)
<code>asffdf_args</code>	further arguments passed to <code>as.ffdf</code> when converting the <code>data.frame</code> of the first chunk to <code>ffdf</code> . Ignored if <code>x</code> is given.
<code>BATCHBYTES</code>	integer: bytes allowed for the size of the <code>data.frame</code> storing the result of reading one chunk. Default <code>getOption("ffbatchbytes")</code> .
<code>VERBOSE</code>	logical: TRUE to verbose timings for each processed chunk (default FALSE)

## Details

`read.table.ffdf` has been designed to read very large (many rows) separated flatfiles in row-chunks and store the result in a `ffdf` object on disk, but quickly accessible via `ff` techniques. The first chunk is read with a default of 1000 rows, for subsequent chunks the number of rows is calculated to not require more RAM than `getOption("ffbatchbytes")`. The following could be indications to change the parameter `first.rows`:

1. set `first.rows=-1` to read the complete file in one go (requires enough RAM)
2. set `first.rows` to a smaller number if the pre-allocation of RAM for the first chunk with parameter `nrows` in `read.table` is too large, i.e. with many columns on machine with little RAM.
3. set `first.rows` to a larger number if you expect better factor level ordering (factor levels are sorted in the first chunk, but not at subsequent chunks, however, factor level ordering can be fixed later, see below).

By default the `ffdf` object is created on the fly at the end of reading the `'first'` chunk, see argument `first.rows`. The creation of the `ffdf` object is done via `as.ffdf` and can be finetuned by passing argument `asffdf_args`. Even more control is possible by passing in a `ffdf` object as argument `x` to which the read records are appended.

`read.table.ffdf` has been designed to behave as much like `read.table` as possible. However, note the following differences:

1. Arguments `'colClasses'` and `'col.names'` are now enforced also during `'next.rows'` chunks. For example giving `colClasses=NA` will force that no `colClasses` are derived from the `first.rows` respective from the `ffdf` object in parameter `x`.
2. `colClass` `'ordered'` is allowed and will create an `ordered` factor
3. character vector are not supported, character data must be read as one of the following `colClasses`: `'Date'`, `'POSIXct'`, `'factor'`, `'ordered'`. By default character columns are read as factors. Accordingly arguments `'as.is'` and `'stringsAsFactors'` are not allowed.
4. the sequence of `levels.ff` from chunked reading can depend on chunk size: by default new levels found on a chunk are appended to the levels found in previous chunks, no attempt is made to sort and recode the levels during chunked processing, levels can be sorted and recoded most efficiently *after* all records have been read using `sortLevels`.
5. the default for argument `'comment.char'` is `""` even for those FUN that have a different default. However, explicit specification of `'comment.char'` will have priority.

### Value

An `ffdf` object. If created during the `'first'` chunk pass, it will have one `physical` component per `virtual` column.

### Note

Note that using the `'skip'` argument still requires to read the file from beginning in order to count the lines to be skipped. If you first read part of the file in order to understand its structure and then want to continue, a more efficient solution that using `'skip'` is opening a `file connection` and pass that to argument `'file'`. `read.table.ffdf` does the same in order to skip efficiently over previously read chunks.

### Author(s)

Jens Oehlschlägel, Christophe Dutang

### See Also

`write.table.ffdf`, `read.table`, `ffdf`

### Examples

```
x <- data.frame(log=rep(c(FALSE, TRUE), length.out=26), int=1:26, db1=1:26 + 0.1
, fac=factor(letters), ord=ordered(LETTERS)
, dct=Sys.time()+1:26, dat=seq(as.Date("1910/1/1"), length.out=26, by=1))
x <- x[c(13:1, 13:1),]
csvfile <- tempPathFile(path=getOption("fftempdir"), extension="csv")
```

```

write.csv(x, file=csvfile, row.names=FALSE)

y <- read.csv(file=csvfile, header=TRUE)
y
cat("Read csv with header\n")
ffx <- read.csv.ffdf(file=csvfile, header=TRUE)
ffx
sapply(ffx[,], class)

message("NOTE that read.table fails for ordered factors, this is fixed in read.table.ffdf")
try(read.csv(file=csvfile, header=TRUE, colClasses=c(ord="ordered")))
# TODO could fix this with the following two commands (Gabor Grothendieck)
# but does not know what bad side-effects this could have
#setOldClass("ordered")
#setAs("character", "ordered", function(from) ordered(from))
y <- read.csv(file=csvfile, header=TRUE, colClasses=c(dct="POSIXct", dat="Date"))
y

ffx <- read.csv.ffdf(file=csvfile, header=TRUE
, colClasses=c(ord="ordered", dct="POSIXct", dat="Date"))
ffx
sapply(ffx[,], class)

message("NOTE that reading in chunks can change the sequence of levels and thus the coding")
message("(Sorting levels during chunked reading can be too expensive)")
ffx <- read.csv.ffdf(file=csvfile, header=TRUE
, colClasses=c(ord="ordered", dct="POSIXct", dat="Date"), first.rows=6, next.rows=10
, VERBOSE=TRUE)
y <- ffx$fac[]
print(levels(y))
data.frame(values=as.character(y), codes=as.integer(y))

message("If we don't know the levels we can sort then after reading")
message("(Will rewrite all factor codes)")
message("NOTE that you MUST assign the return value of sortLevels()")
ffx <- sortLevels(ffx)
y <- ffx$fac[]
print(levels(y))
data.frame(values=as.character(y), codes=as.integer(y))

message("If we KNOW the levels we can fix levels upfront")
ffx <- read.csv.ffdf(file=csvfile, header=TRUE
, colClasses=c(ord="ordered", dct="POSIXct", dat="Date")
, first.rows=6, next.rows=10, levels=list(fac=letters, ord=LETTERS))
y <- ffx$fac[]
print(levels(y))
data.frame(values=as.character(y), codes=as.integer(y))

message("Or we inspect a sufficiently large chunk of data and use those")
ffx1 <- read.csv.ffdf(file=csvfile, header=TRUE
, colClasses=c(ord="ordered", dct="POSIXct", dat="Date"), nrows=13)
ffx <- read.csv.ffdf(x=ffx1, file=csvfile, header=FALSE, skip=1+nrow(ffx1), VERBOSE=TRUE)

```

```

    message("We can check for unexpected factor levels, say we only allowed a:1")
    ffx <- read.csv.ffdf(file=csvfile, header=TRUE
, colClasses=c(ord="ordered", dct="POSIXct", dat="Date")
, levels=list(fac=letters[1:12], ord=LETTERS[1:12]), appendLevels=FALSE)
    sapply(colnames(ffx), function(i)sum(is.na(ffx[[i]][,])))

message("We can fine-tune the creation of the ffdf:")
message("- let's create the ff files outside of fftempdir")
message("- let's reduce required disk space and thus file.system cache RAM")
vmode(ffx)
message("By default we had record size")
sum(.ffbytes[vmode(ffx)])

ffx <- read.csv.ffdf(file=csvfile, header=TRUE
, colClasses=c(ord="ordered", dct="POSIXct", dat="Date")
, asffdf_args=list(
  vmode = c(log="boolean", int="byte", dbl="single", fac="nibble"
, ord="nibble", dct="single", dat="single")
, col_args=list(pattern = "./csv") # create in getwd() with prefix csv
)
)
vmode(ffy)
message("This recordsize is more than 50% reduced")
sum(.ffbytes[vmode(ffy)])

message("Don't forget to wrap-up files that are not in fftempdir")
delete(ffy); rm(ffy)
message("It's a good habit to also wrap-up temporary stuff (or at least know how this is done)")
rm(ffx); gc()

fwffile <- tempfile()

cat(file=fwffile, "123456", "987654", sep="\n")
x <- read.fwf(fwffile, widths=c(1,2,3)) #> 1 23 456 \ 9 87 654
y <- read.table.ffdf(file=fwffile, FUN="read.fwf", widths=c(1,2,3))
stopifnot(identical(x, y[,]))
x <- read.fwf(fwffile, widths=c(1,-2,3)) #> 1 456 \ 9 654
y <- read.table.ffdf(file=fwffile, FUN="read.fwf", widths=c(1,-2,3))
stopifnot(identical(x, y[,]))
unlink(fwffile)

cat(file=fwffile, "123", "987654", sep="\n")
x <- read.fwf(fwffile, widths=c(1,0, 2,3)) #> 1 NA 23 NA \ 9 NA 87 654
y <- read.table.ffdf(file=fwffile, FUN="read.fwf", widths=c(1,0, 2,3))
stopifnot(identical(x, y[,]))
unlink(fwffile)

cat(file=fwffile, "123456", "987654", sep="\n")
x <- read.fwf(fwffile, widths=list(c(1,0, 2,3), c(2,2,2))) #> 1 NA 23 456 98 76 54
y <- read.table.ffdf(file=fwffile, FUN="read.fwf", widths=list(c(1,0, 2,3), c(2,2,2)))
stopifnot(identical(x, y[,]))
unlink(fwffile)

```

```
unlink(csvfile)
```

---

```
readwrite.ff
```

```
Reading and writing vectors (low-level)
```

---

## Description

Simple low-level interface for reading and writing vectors from ff files.

## Usage

```
read.ff(x, i, n)
write.ff(x, i, value, add = FALSE)
readwrite.ff(x, i, value, add = FALSE)
```

## Arguments

x	an ff object
i	a start position in the ff file
n	number of elements to read
value	vector of elements to write
add	TRUE if the values should rather increment than overwrite at the target positions

## Details

`readwrite.ff` combines the effects of `read.ff` and `write.ff` in a single operation: it retrieves the old values starting from position `i` before changing them. `getset.ff` will maintain `na.count`.

## Value

`read.ff` returns a vector of values, `write.ff` returns the 'changed' ff object (like all assignment functions do) and `readwrite.ff` returns the values at the target position. More precisely `readwrite.ff(x, i, value, add=FALSE)` returns the old values at the position `i` while `readwrite.ff(x, i, value, add=TRUE)` returns the incremented values of `x`.

## Note

`read.ff`, `write.ff` and `readwrite.ff` are low level functions that do not support `ramclass` and `ramattribs` and thus will not give the expected result with `factor` and `POSIXct`

## Author(s)

Jens Oehlschlägel

**See Also**

[getset.ff](#) for low-level scalar access and [\[.ff](#) for high-level access

**Examples**

```
x <- ff(0, length=12)
read.ff(x, 3, 6)
write.ff(x, 3, rep(1, 6))
x
write.ff(x, 3, rep(1, 6), add=TRUE)
x
readwrite.ff(x, 3, rep(1, 6), add=TRUE)
readwrite.ff(x, 3, rep(1, 6))
x
rm(x); gc()
```

---

regtest.fforder

*Sorting: regression tests*

---

**Description**

Some tests verifying the correctness of the sorting routines

**Usage**

```
regtest.fforder(n = 100)
```

**Arguments**

n                    size of vector to be sorted

**Details**

stops in case of an error

**Value**

Invisible()

**Author(s)**

Jens Oehlschlägel

**See Also**

[ramsort](#)

**Examples**

```

regtest.fforder()

## Not run:
n <- 5e6
message("performance comparison at n=", n, "")

message("sorting doubles")
x <- y <- as.double(runif(n))

x[] <- y
system.time(sort(x))[3]
x[] <- y
system.time(shellsort(x))[3]
x[] <- y
system.time(shellsort(x, has.na=FALSE))[3]
x[] <- y
system.time(mergesort(x))[3]
x[] <- y
system.time(mergesort(x, has.na=FALSE))[3]

x[] <- y
system.time(sort(x, decreasing=TRUE))[3]
x[] <- y
system.time(shellsort(x, decreasing=TRUE))[3]
x[] <- y
system.time(shellsort(x, decreasing=TRUE, has.na=FALSE))[3]
x[] <- y
system.time(mergesort(x, decreasing=TRUE))[3]
x[] <- y
system.time(mergesort(x, decreasing=TRUE, has.na=FALSE))[3]

x <- y <- as.double(sample(c(rep(NA, n/2), runif(n/2))))

x[] <- y
system.time(sort(x))[3]
x[] <- y
system.time(shellsort(x))[3]
x[] <- y
system.time(mergesort(x))[3]

x[] <- y
system.time(sort(x, decreasing=TRUE))[3]
x[] <- y
system.time(shellsort(x, decreasing=TRUE))[3]
x[] <- y
system.time(mergesort(x, decreasing=TRUE))[3]

x <- y <- sort(as.double(runif(n)))

```



```
x[] <- y
system.time(sort(x)) # only here R is faster because R checks for being sorted
x[] <- y
system.time(shellsort(x))[3]
x[] <- y
system.time(shellsort(x, has.na=FALSE))[3]
x[] <- y
system.time(mergesort(x))[3]
x[] <- y
system.time(mergesort(x, has.na=FALSE))[3]

x[] <- y
system.time(sort(x, decreasing=TRUE))[3]
x[] <- y
system.time(shellsort(x, decreasing=TRUE))[3]
x[] <- y
system.time(shellsort(x, decreasing=TRUE, has.na=FALSE))[3]
x[] <- y
system.time(mergesort(x, decreasing=TRUE))[3]
x[] <- y
system.time(mergesort(x, decreasing=TRUE, has.na=FALSE))[3]

y <- rev(y)
x[] <- y
system.time(sort(x))[3]
x[] <- y
system.time(shellsort(x))[3]
x[] <- y
system.time(shellsort(x, has.na=FALSE))[3]
x[] <- y
system.time(mergesort(x))[3]
x[] <- y
system.time(mergesort(x, has.na=FALSE))[3]

x[] <- y
system.time(sort(x, decreasing=TRUE))[3]
x[] <- y
system.time(shellsort(x, decreasing=TRUE))[3]
x[] <- y
system.time(shellsort(x, decreasing=TRUE, has.na=FALSE))[3]
x[] <- y
system.time(mergesort(x, decreasing=TRUE))[3]
x[] <- y
system.time(mergesort(x, decreasing=TRUE, has.na=FALSE))[3]

rm(x,y)

message("ordering doubles")

x <- as.double(runif(n))
system.time(order(x))[3]
```

```
i <- 1:n
system.time(shellorder(x, i))[3]
i <- 1:n
system.time(shellorder(x, i, stabilize=TRUE))[3]
i <- 1:n
system.time(mergeorder(x, i))[3]

x <- as.double(sample(c(rep(NA, n/2), runif(n/2))))
system.time(order(x))[3]
i <- 1:n
system.time(shellorder(x, i))[3]
i <- 1:n
system.time(shellorder(x, i, stabilize=TRUE))[3]
i <- 1:n
system.time(mergeorder(x, i))[3]

x <- as.double(sort(runif(n)))
system.time(order(x))[3]
i <- 1:n
system.time(shellorder(x, i))[3]
i <- 1:n
system.time(shellorder(x, i, stabilize=TRUE))[3]
i <- 1:n
system.time(mergeorder(x, i))[3]

x <- rev(x)
system.time(order(x))[3]
i <- 1:n
system.time(shellorder(x, i))[3]
i <- 1:n
system.time(shellorder(x, i, stabilize=TRUE))[3]
i <- 1:n
system.time(mergeorder(x, i))[3]

x <- as.double(runif(n))
system.time(order(x, decreasing=TRUE))[3]
i <- 1:n
system.time(shellorder(x, i, decreasing=TRUE))[3]
i <- 1:n
system.time(shellorder(x, i, decreasing=TRUE, stabilize=TRUE))[3]
i <- 1:n
system.time(mergeorder(x, i, decreasing=TRUE))[3]

x <- as.double(sample(c(rep(NA, n/2), runif(n/2))))
system.time(order(x, decreasing=TRUE))[3]
i <- 1:n
system.time(shellorder(x, i, decreasing=TRUE))[3]
i <- 1:n
system.time(shellorder(x, i, decreasing=TRUE, stabilize=TRUE))[3]
i <- 1:n
system.time(mergeorder(x, i, decreasing=TRUE))[3]
```

```

x <- as.double(sort(runif(n)))
system.time(order(x, decreasing=TRUE))[3]
i <- 1:n
system.time(shellorder(x, i, decreasing=TRUE))[3]
i <- 1:n
system.time(shellorder(x, i, decreasing=TRUE, stabilize=TRUE))[3]
i <- 1:n
system.time(mergeorder(x, i, decreasing=TRUE))[3]

x <- rev(x)
system.time(order(x, decreasing=TRUE))[3]
i <- 1:n
system.time(shellorder(x, i, decreasing=TRUE))[3]
i <- 1:n
system.time(shellorder(x, i, decreasing=TRUE, stabilize=TRUE))[3]
i <- 1:n
system.time(mergeorder(x, i, decreasing=TRUE))[3]

keys <- c("short", "ushort")
for (v in c("integer", keys)){

  if (v %in% keys){
    k <- .vmax[v]-.vmin[v]+1L
    if (is.na(.vNA[v])){
      y <- sample(c(rep(NA, k), .vmin[v]:.vmax[v]), n, TRUE)
    }else{
      y <- sample(.vmin[v]:.vmax[v], n, TRUE)
    }
  }else{
    k <- .Machine$integer.max
    y <- sample(k, n, TRUE)
  }

  message("sorting ", v)

  x <- y
  message("sort(x) ", system.time(sort(x))[3])
  x <- y
  message("shellsort(x) ", system.time(shellsort(x))[3])
  x <- y
  message("mergesort(x) ", system.time(mergesort(x))[3])
  x <- y
  message("radixsort(x) ", system.time(radixsort(x))[3])
  if (v %in% keys){
    x <- y
    message("keysort(x) ", system.time(keysort(x))[3])
    x <- y
    message("keysort(x, keyrange=c(.vmin[v],.vmax[v])) "
, system.time(keysort(x, keyrange=c(.vmin[v],.vmax[v])))[3])
  }

  if (!is.na(.vNA[v])){

```

```

x <- y
message("shellsort(x, has.na=FALSE) ", system.time(shellsort(x, has.na=FALSE))[3])
x <- y
message("mergesort(x, has.na=FALSE) ", system.time(mergesort(x, has.na=FALSE))[3])
x <- y
message("radixsort(x, has.na=FALSE) ", system.time(radixsort(x, has.na=FALSE))[3])
if (v %in% keys){
  x <- y
  message("keysort(x, has.na=FALSE) ", system.time(keysort(x, has.na=FALSE))[3])
  x <- y
  message("keysort(x, has.na=FALSE, keyrange=c(.vmin[v],.vmax[v])) "
, system.time(keysort(x, has.na=FALSE, keyrange=c(.vmin[v],.vmax[v])))[3])
}
}

message("ordering",v)

x[] <- y
i <- 1:n
message("order(x) ", system.time(order(x))[3])
x[] <- y
i <- 1:n
message("shellorder(x, i) ", system.time(shellorder(x, i))[3])
x[] <- y
i <- 1:n
message("mergeorder(x, i) ", system.time(mergeorder(x, i))[3])
x[] <- y
i <- 1:n
message("radixorder(x, i) ", system.time(radixorder(x, i))[3])
if (v %in% keys){
  x[] <- y
  i <- 1:n
  message("keyorder(x, i) ", system.time(keyorder(x, i))[3])
  x[] <- y
  i <- 1:n
  message("keyorder(x, i, keyrange=c(.vmin[v],.vmax[v])) "
, system.time(keyorder(x, i, keyrange=c(.vmin[v],.vmax[v])))[3])
}

if (!is.na(.vNA[v])){
  x[] <- y
  i <- 1:n
  message("shellorder(x, i, has.na=FALSE) ", system.time(shellorder(x, i, has.na=FALSE))[3])
  x[] <- y
  i <- 1:n
  message("mergeorder(x, i, has.na=FALSE) ", system.time(mergeorder(x, i, has.na=FALSE))[3])
  x[] <- y
  i <- 1:n
  message("radixorder(x, i, has.na=FALSE) ", system.time(radixorder(x, i, has.na=FALSE))[3])
  if (v %in% keys){
    x[] <- y
    i <- 1:n

```

```

        message("keyorder(x, i, has.na=FALSE) ", system.time(keyorder(x, i, has.na=FALSE))[3])
        x[] <- y
        i <- 1:n
        message("keyorder(x, i, has.na=FALSE, keyrange=c(.vmin[v],.vmax[v])) "
, system.time(keyorder(x, i, has.na=FALSE, keyrange=c(.vmin[v],.vmax[v])))[3])
    }
}

}

## End(Not run)

```

---

repnam

*Replicate with names*


---

### Description

Function `repnam` replicates its argument to the desired length, either by simply [replicating](#) or - if it has [names](#) - by replicating the default and matching the argument by its names.

### Usage

```
repnam(argument, names = NULL, len=length(names), default = list(NULL))
```

### Arguments

<code>argument</code>	a named or non-named vector or list to be replicated
<code>names</code>	NULL or a character vector of names to which the argument names are matched
<code>len</code>	the desired length (required if <code>names</code> is not given)
<code>default</code>	the desired default which is replicated in case names are used (the default <code>list(NULL)</code> is suitable for a list argument)

### Value

an object like `argument` or `default` having length `len`

### Note

This is for internal use, e.g. to handle argument `colClasses` in [read.table.ffdf](#)

### Author(s)

Jens Oehlschlägel

### See Also

[rep](#), [vector](#), [repfromto](#)

**Examples**

```

message("a list example")
replam(list(y=c(1,2), z=3), letters)
replam(list(c(1,2), 3), letters)

message("a vector example")
replam(c(y=1, z=3), letters, default=NA)
replam(c(1, 3), letters, default=NA)

```

---

sortLevels

*Factor level manipulation*


---

**Description**

appendLevels combines [levels](#) without sorting such that levels of the first argument will not require re-coding.

recodeLevels is a generic for recoding a factor to a desired set of levels - also has a method for large [ff](#) objects

sortLevels is a generic for level sorting and recoding of single factors or of all factors of a [ffdf](#) dataframe.

**Usage**

```

appendLevels(...)
recodeLevels(x, lev)
## S3 method for class 'factor'
recodeLevels(x, lev)
## S3 method for class 'ff'
recodeLevels(x, lev)
sortLevels(x)
## S3 method for class 'factor'
sortLevels(x)
## S3 method for class 'ff'
sortLevels(x)
## S3 method for class 'ffdf'
sortLevels(x)

```

**Arguments**

...	character vector of levels or <a href="#">is.factor</a> objects from which the level attribute is taken
x	a <a href="#">factor</a> or <a href="#">ff</a> factor or a <a href="#">ffdf</a> dataframe (sortLevels only)
lev	a character vector of levels

**Details**

When reading a long file with categorical columns the final set of factor levels is only known once the complete file has been read. When a file is so large that we read it in chunks, the new levels need to be added incrementally. `rbind.data.frame` sorts combined levels, which requires recoding. For `ff` factors this would require recoding of all previous chunks at the next chunk - potentially on disk, which is too expensive. Therefore `read.table.ffdf` will simply `appendLevels` without sorting, and the `recodeLevels` and `sortLevels` generics provide a convenient means for sorting and recoding levels after all chunks have been read.

**Value**

`appendLevels` returns a vector of combined levels, `recodeLevels` and `sortLevels` return the input object with changed levels. Do read the note!

**Note**

You need to re-assign the return value not only for `ram`- but also for `ff`-objects. Remember `ff`'s hybrid copying semantics: `LimWarn`. If you forget to re-assign the returned object, you will end up with `ff` objects that have their integer codes re-coded to the new levels but still carry the old levels as a `virtual` attribute.

**Author(s)**

Jens Oehlschlägel

**See Also**

[read.table.ffdf](#), [levels.ff](#)

**Examples**

```
message("Let's create a factor with little levels")
x <- ff(letters[4:6], levels=letters[4:6])
message("Let's interpret the same ff file without levels in order to see the codes")
y <- x
levels(y) <- NULL

levels(x)
data.frame(factor=x[], codes=y[])

levels(x) <- appendLevels(levels(x), letters)
levels(x)
data.frame(factor=x[], codes=y[])

x <- sortLevels(x) # implicit recoding is chunked were necessary
levels(x)
data.frame(factor=x[], codes=y[])

message("NEVER forget to reassign the result of recodeLevels or sortLevels,
look at the following mess")
recodeLevels(x, rev(levels(x)))
```

```

message("NOW the codings have changed, but not the levels, the result is wrong data")
levels(x)
data.frame(factor=x[], codes=y[])

rm(x);gc()

## Not run:
n <- 5e7

message("reading a factor from a file ist as fast ...")
system.time(
fx <- ff(factor(letters[1:25]), length=n)
)
system.time(x <- fx[])
str(x)
rm(x); gc()

message("... as creating it in-RAM (R-2.11.1) which is theoretically impossible ...")
system.time({
x <- integer(n)
x[] <- 1:25
levels(x) <- letters[1:25]
class(x) <- "factor"
})
str(x)
rm(x); gc()

message("... but is possible if we avoid some unnecessary copying that is triggered
by assignment functions")
system.time({
x <- integer(n)
x[] <- 1:25
setattr(x, "levels", letters[1:25])
setattr(x, "class", "factor")
})
str(x)
rm(x); gc()

rm(n)

## End(Not run)

```

---

splitPathFile

*Analyze pathfile-strings*


---

### Description

splitPathFile splits a vector of pathfile-strings into path- and file-components without loss of



information. `unsplitPathFile` restores the original pathfile-string vector. `standardPathFile` standardizes a vector of pathfile-strings: backslashes are replaced by slashes, except for the first two leading backslashes indicating a network share. `tempPathFile` returns - similar to `tempfile` - a vector of filenames given path(s) and file-prefix(es) and an optional extension. `fftempfile` returns - similar to `tempPathFile` - a vector of filenames following a vector of pathfile patterns that are interpreted in a ff-specific way.

## Usage

```
splitPathFile(x)
unsplitPathFile(splitted)
standardPathFile(x)
tempPathFile(splitted=NULL, path=splitted$path, prefix=splitted$file, extension=NULL)
fftempfile(x)
```

## Arguments

<code>x</code>	a character vector of pathfile strings
<code>splitted</code>	a return value from <code>splitPathFile</code>
<code>path</code>	a character vector of path components
<code>prefix</code>	a character vector of file components
<code>extension</code>	optional extension like "csv" (or NULL)

## Details

`dirname` and `basename` remove trailing file separators and therefore cannot distinguish pathfile string that contains ONLY a path from a pathfile string that contains a path AND file. Therefore `file.path(dirname(pathfile), basename(pathfile))` cannot always restore the original pathfile string.

`splitPathFile` decomposes each pathfile string into three parts: a path BEFORE the last file separator, the file separator, the filename component AFTER the last file separator. If there is no file separator in the string, `splitPathFile` tries to guess whether the string is a path or a file component: ".", ".." and "~" are recognized as path components. No tilde expansion is done, see `path.expand`. Backslashes are converted to the current `.Platform$file.sep` using `splitPathFile` except for the first two leading backslashes indicating a network share.

`unsplitPathFile` restores the original pathfile-string vector up to translated backslashes.

`tempPathFile` internally uses `tempfile` to create its filenames, if an extension is given it repeats filename creation until none of them corresponds to an existing file.

`fftempfile` takes a path-prefix pattern as input, splits it, will replace an empty path by `getOption("fftempdir")` and will use `getOption("ffextension")` as extension.

## Value

A list with components

<code>path</code>	a character vector of path components
<code>fsep</code>	a character vector of file separators or ""
<code>file</code>	a character vector of file components

**Note**

There is no guarantee that the path and file components contain valid path- or file-names. Like [basename](#), `splitPathFile` can return ".", ".." or even "", however, all these make sense as a prefix in `tempPathFile`.

**Author(s)**

Jens Oehlschlägel

**See Also**

[tempfile](#), [dirname](#), [basename](#), [file.path](#)

**Examples**

```

pathfile <- c("", ".", "/.", "./", "./", "/",
, "a", "a/", "/a", "a/a", ".a", "a/.", "c:/a/b/c", "c:/a/b/c/"
, "..", "../", "/.", "../.", "//", "\\a\\", "\\a/"
, "\\a/b", "\\a/b/", "~", "~/", "~/a", "~/a/")
splitted <- splitPathFile(pathfile)
restored <- unsplitPathFile(splitted)
stopifnot(all(gsub("\\", "/", restored) == gsub("\\", "/", pathfile)))

dirnam <- dirname(pathfile)
basnam <- basename(pathfile)

db <- file.path(dirnam, basnam)
ident = gsub("\\", "/", db) == gsub("\\", "/", pathfile)
sum(!ident)

do.call("data.frame", c(list(ident=ident, pathfile=pathfile
, dirnam=dirnam, basnam=basnam), splitted))

## Not run:
message("show the difference between tempfile and ftempfile")
do.call("data.frame", c(list(ident=ident, pathfile=pathfile, dirnam=dirnam, basnam=basnam)
, splitted, list(filename=tempPathFile(splitted), ftempfile=ftempfile(pathfile))))

message("for a single string splitPathFile is slower,
for vectors of strings it scales much better than dirname+basename")

system.time(for (i in 1:1000){
  d <- dirname(pathfile)
  b <- basename(pathfile)
})
system.time(for (i in 1:1000){
  s <- splitPathFile(pathfile)
})

len <- c(1,10,100,1000)
timings <- matrix(0, 2, length(len), dimnames=list(c("dir.base.name", "splitPathFile"), len))
for (j in seq(along=len)){

```

```

l <- len[j]
r <- 10000 / l
x <- rep("\\\\a/b/", l)
timings[1,j] <- system.time(for (i in 1:r){
  d <- dirname(x)
  b <- basename(x)
})[3]
timings[2,j] <- system.time(for (i in 1:r){
  s <- splitPathFile(x)
})[3]
}
timings

## End(Not run)

```

---

swap

*Reading and writing in one operation (high-level)*


---

## Description

The generic swap combines `x[i]` and `x[i] <- value` in a single operation.

## Usage

```

swap(x, value, ...)
## S3 method for class 'ff'
swap(x, value, i, add = FALSE, pack = FALSE, ...)
## S3 method for class 'ff_array'
swap(x, value, ..., bydim = NULL, drop = getOption("ffdrop"), add = FALSE, pack = FALSE)
## Default S3 method:
swap(x, value, ..., add = FALSE)

```

## Arguments

<code>x</code>	a ff or ram object
<code>value</code>	the new values to write, possibly recycled, see <a href="#">[.ff]</a>
<code>i</code>	index information, see <a href="#">[.ff]</a>
<code>...</code>	missing OR up to <code>length(dim(x))</code> index expressions OR (ff only) <a href="#">hi</a> objects
<code>drop</code>	logical scalar indicating whether array dimensions shall be dropped
<code>bydim</code>	how to interpret vector to array data, see <a href="#">[.ff]</a>
<code>add</code>	TRUE if the values should rather increment than overwrite at the target positions, see <a href="#">readwrite.ff</a>
<code>pack</code>	FALSE to prevent rle-packing in hybrid index preprocessing, see <a href="#">as.hi</a>

**Details**

```
y <- swap(x, value, i, add=FALSE, ...)
```

is a shorter and more efficient version of

```
y <- x[i, add=FALSE, ...]
x[i, add=FALSE, ...] <- value
```

and

```
y <- swap(x, value, i, add=TRUE, ...)
```

is a shorter and more efficient version of

```
y <- x[i, add=TRUE, ...]
y <- y + value
x[i, add=FALSE, ...] <- y
```

**Value**

Values at the target positions. More precisely `swap(x, value, i, add=FALSE)` returns the old values at the position `i` while `swap(x, value, i, add=TRUE)` returns the incremented values of `x`.

**Note**

Note that `swap.default` changes the object in its parent frame and thus violates R's usual functional programming logic. When using `add=TRUE`, duplicated index positions should be avoided, because `ff` and `ram` objects behave differently:

```
swap.ff(x, 1, c(3,3), add=TRUE)
# will increment x at position 3 TWICE by 1, while
swap.default(x, 1, c(3,3), add=TRUE)
# will increment x at position 3 just ONCE by 1
```

**Author(s)**

Jens Oehlschlägel

**See Also**

[\[.ff, add, readwrite.ff, getset.ff, LimWarn](#)

**Examples**

```
x <- ff("a", levels=letters, length=52)
y <- swap(x, "b", sample(length(x), 26))
```

```
x  
y  
rm(x,y); gc()
```

---

symmetric

*Test for symmetric structure*

---

## Description

Check if an object is inherently symmetric (its structure, not its data)

## Usage

```
symmetric(x, ...)  
## S3 method for class 'ff'  
symmetric(x, ...)  
## Default S3 method:  
symmetric(x, ...)  
## S3 method for class 'dist'  
symmetric(x, ...)
```

## Arguments

x	an ff or ram object
...	further arguments (not used)

## Details

ff matrices can be declared symmetric at creation time. Compatibility function `symmetric.default` returns FALSE, `symmetric.dist` returns TRUE.

## Value

TRUE or FALSE

## Author(s)

Jens Oehlschlägel

## See Also

[symmetric](#), [ff](#), [dist](#), [isSymmetric](#)

## Examples

```
symmetric(matrix(1:16, 4, 4))  
symmetric(dist(rnorm(1:4)))
```

---

symmIndex2vectorIndex *Array: make vector positions from symmetric array index*

---

### Description

make vector positions from (non-symmetric) array index respecting 'dim' and 'fixdiag'

### Usage

```
symmIndex2vectorIndex(x, dim, fixdiag = NULL)
```

### Arguments

x	a matrix[,1:2] with matrix subscripts
dim	the dimensions of the symmetric matrix
fixdiag	NULL assumes free diagonal, any value assumes fixed diagonal

### Details

With 'fixdiag = NULL'

### Value

a vector of indices in 1:prod(dim(x))

### Author(s)

Jens Oehlschlägel

### See Also

[arrayIndex2vectorIndex](#)

### Examples

```
symmIndex2vectorIndex(rbind(
  c(1,1)
  ,c(1,10)
  ,c(10,1)
  ,c(10,10)
), dim=c(10,10))
symmIndex2vectorIndex(rbind(
  c(1,1)
  ,c(1,10)
  ,c(10,1)
  ,c(10,10)
), dim=c(10,10), fixdiag=1)
```

---

unclass_-	<i>Unclassed assignement</i>
-----------	------------------------------

---

**Description**

With `unclass<-` you can circumvent class dispatch on the assignment operator

**Usage**

```
unclass(x) <- value
```

**Arguments**

x	some object
value	the value to be assigned

**Value**

the modified object

**Author(s)**

Jens Oehlschlägel

**See Also**

[unclass](#), [undim](#)

**Examples**

```
x <- factor(letters)
unclass(x)[1:3] <- 1L
x
```

---

undim	<i>Undim</i>
-------	--------------

---

**Description**

`undim` returns its input with the `dim` attribute removed

**Usage**

```
undim(x)
```

**Arguments**

x                    an object

**Value**

x without dim attribute

**Author(s)**

Jens Oehlschlägel

**See Also**

[unclass<-](#), [unclass](#), [unname](#), [dim](#)

**Examples**

```
x <- matrix(1:12, 3)
x
undim(x)
```

---

unsort

*Hybrid Index, internal utilities*

---

**Description**

Non-documented internal utilities that might change

**Usage**

```
unsort(x, ix)
unsort.hi(x, index)
unsort.ahi(x, index, ixre = any(sapply(index, function(i) {
  if (is.null(i$ix)) {
    if (i$re) TRUE else FALSE
  } else {
    TRUE
  }
})), ix = lapply(index, function(i) {
  if (is.null(i$ix)) {
    if (i$re)
      orig <- rev(1:poslength(i))
    else orig <- 1:poslength(i)
  }
  else {
    orig <- i$ix
  }
})
```



```

    orig
  )))
  subscript2integer(x, maxindex = NULL, names = NULL)

```

### Arguments

x	x
ix	ix
ixre	ixre
index	index
maxindex	maxindex
names	names

### Details

These are utility functions for restoring original order after sorting. For now we 'mimic' the intuitive but wrong argument order of `match()` which should rather have the 'table' argument as its first argument, then one could properly method-dispatch on the type of table. xx We might change to proper 'unsort' generic, but then we have to change argument order.

### Value

undefined

### Author(s)

Jens Oehlschlägel

### See Also

[hi](#), [as.hi](#)

---

update.ff

*Update ff content from another object*

---

### Description

update copies updates one ff object with the content of another object.

### Usage

```

## S3 method for class 'ff'
update(object, from, delete = FALSE, bydim = NULL, fromdim = NULL
, BATCHSIZE = .Machine$integer.max, BATCHBYTES = getOption("ffbatchbytes")
, VERBOSE = FALSE, ...)
## S3 method for class 'ffdf'
update(object, from, ...)

```

**Arguments**

object	an ff object to which to update
from	an object from which to uodate
delete	NA for quick update with file-exchange, TRUE for quick update with deleting the 'from' object after the update, can speed up updating significantly
bydim	how to interpret the content of the object, see <a href="#">ff</a>
fromdim	how to interpret the content of the 'from' object, see <a href="#">ff</a>
BATCHSIZE	BATCHSIZE
BATCHBYTES	BATCHBYTES
VERBOSE	VERBOSE
...	further arguments

**Details**

If the source object is `ff` and not `delete=FALSE` then instead of slow copying we - if possible - try to swap and rename the files behind the `ff` objects. Quick update requires that the two `ff` objects are [vectorCompatible](#), that both don't use [vw](#), that they have identical [maxlength](#) and identical [levels.ff](#).

**Value**

An `ff` object like the input 'object' updated with the content of the 'from' object.

**Note**

You don't have a guarantee that with `delete=TRUE` the 'from' object gets deleted or with `delete=NA` the 'from' objects carries the content of 'object'. Such expectations only turn true if really a quick update was possible.

**Author(s)**

Jens Oehlschlägel

**See Also**

[ff](#), [clone](#), [ffvecapply](#), [vectorCompatible](#), [filename](#)

**Examples**

```
x <- ff(1:100)
y <- ff(-(1:100))
message("You should make it a habit to re-assign the return value
of update although this is not needed currently.")
x <- update(x, from=y)
x
y
x[] <- 1:100
```

```

x <- update(x, from=y, delete=NA)
x
y
x <- update(x, from=y, delete=TRUE)
x
y
x
rm(x,y); gc()

## Not run:
message("timings")
x <- ff(1:1000000)
y <- ff(-(1:1000000))
system.time(update(x, from=y))
system.time(update(y, from=x, delete=NA))
system.time(update(x, from=y, delete=TRUE))
rm(x,y); gc()

## End(Not run)

```

---

vecprint

*Print beginning and end of big vector*


---

## Description

Print beginning and end of big vector

## Usage

```

vecprint(x, maxlength = 16, digits = getOption("digits"))
## S3 method for class 'vecprint'
print(x, quote = FALSE, ...)

```

## Arguments

x	a vector
maxlength	max number of elements for printing
digits	see <a href="#">format</a>
quote	see <a href="#">print</a>
...	see <a href="#">print</a>

## Value

a list of class 'vecprint' with components

subscript	a list with two vectors of subscripts: vector begin and vector end
example	the extracted example vector as.character including separator
sep	the row separator ":"

**Author(s)**

Jens Oehlschlägel

**See Also**[matprint](#)**Examples**

```
vecprint(10000:1)
```

---

`vector.vmode`*Create vector of virtual mode*

---

**Description**

`vector.vmode` creates a vector of a given `vmode` and length

**Usage**

```
vector.vmode(vmode = "logical", length = 0)
boolean(length = 0)
quad(length = 0)
nibble(length = 0)
byte(length = 0)
ubyte(length = 0)
short(length = 0)
ushort(length = 0)
```

**Arguments**

<code>vmode</code>	virtual mode
<code>length</code>	desired length

**Details**

Function `vector.vmode` creates the vector in one of the usual [storage.modes](#) (see [.rammode](#)) but flags them with an additional attribute 'vmode' if necessary. The creators can also be used directly:

<code>boolean</code>	1 bit logical without NA
<code>logical</code>	2 bit logical with NA
<code>quad</code>	2 bit unsigned integer without NA
<code>nibble</code>	4 bit unsigned integer without NA
<code>byte</code>	8 bit signed integer with NA
<code>ubyte</code>	8 bit unsigned integer without NA
<code>short</code>	16 bit signed integer with NA
<code>ushort</code>	16 bit unsigned integer without NA

integer	32 bit signed integer with NA
single	32 bit float
double	64 bit float
complex	2x64 bit float
raw	8 bit unsigned char
character	character

### Value

a vector of the desired vmode initialized with 0

### Author(s)

Jens Oehlschlägel

### See Also

[as.vmode](#), [vector](#)

### Examples

```
vector.vmode("byte",12)
vector.vmode("double",12)
byte(12)
double(12)
```

---

vector2array

*Array: make array from vector*

---

### Description

makes array from vector respecting 'dim' and 'dimorder'

### Usage

```
vector2array(x, dim, dimorder = NULL)
```

### Arguments

x	an input vector, recycled if needed
dim	<a href="#">dim</a>
dimorder	<a href="#">dimorder</a>

**Details**

FILLS vector into array of dim where fastest rotating is dim[dimorder[1]], next is dim[dimorder[2]] and so forth. This is a generalization of converting vector to matrix(, byrow=TRUE). NOTE that the result is a ram array always stored in STANDARD dimorder !!! In this usage we sometimes term the dimorder 'bydim' because it does not change the physical layout of the result, rather bydim refers to the dimorder in which to interpret the vector (not the result). In ff, update and clone we have 'bydim' to contrast it from 'dimorder', the latter describing the layout of the file.

**Value**

a suitable [array](#)

**Author(s)**

Jens Oehlschlägel

**See Also**

[array2vector](#), [vectorIndex2arrayIndex](#)

**Examples**

```
vector2array(1:12, dim=c(3, 4))           # matrix(1:12, 3, 4)
vector2array(1:12, dim=c(3, 4), dimorder=2:1) # matrix(1:12, 3, 4, byrow=TRUE)
```

---

vectorIndex2arrayIndex

*Array: make array from index vector positions*

---

**Description**

make array from index vector positions respecting 'dim' and 'dimorder'

**Usage**

```
vectorIndex2arrayIndex(x, dim = NULL, dimorder = NULL, vw = NULL)
```

**Arguments**

x	a vector of indices in 1:prod(dim)
dim	NULL or <a href="#">dim</a>
dimorder	NULL or <a href="#">dimorder</a>
vw	NULL or integer matrix[2,m], see details

**Details**

The fastest rotating dimension is `dim[dimorder[1]]`, then `dim[dimorder[2]]`, and so forth.

The parameters 'x' and 'dim' may refer to a subarray of a larger array, in this case, the array indices 'x' are interpreted as 'vw[1,] + x' within the larger array 'vw[1,] + x + vw[2,]'.

**Value**

an n by m matrix with n m-dimensional array indices

**Author(s)**

Jens Oehlschlägel

**See Also**

[vector2array](#), [arrayIndex2vectorIndex](#), [symmIndex2vectorIndex](#)

**Examples**

```
matrix(1:12, 3, 4)
vectorIndex2arrayIndex(1:12, dim=3:4)
vectorIndex2arrayIndex(1:12, dim=3:4, dimorder=2:1)
matrix(1:30, 5, 6)
vectorIndex2arrayIndex(c(6L, 7L, 8L, 11L, 12L, 13L, 16L, 17L, 18L, 21L, 22L, 23L)
, vw=rbind(c(0,1), c(3,4), c(2,1)))
vectorIndex2arrayIndex(c(2L, 8L, 14L, 3L, 9L, 15L, 4L, 10L, 16L, 5L, 11L, 17L)
, vw=rbind(c(0,1), c(3,4), c(2,1)), dimorder=2:1)
```

---

vmode

*Virtual storage mode*

---

**Description**

Function `vmode` returns virtual storage modes of 'ram' or 'ff' objects, the generic `vmode<-` sets the `vmode` of ram objects (`vmode` of ff objects cannot be changed).

**Usage**

```
vmode(x, ...)
vmode(x) <- value
## Default S3 method:
vmode(x, ...)
## S3 method for class 'ff'
vmode(x, ...)
## Default S3 replacement method:
vmode(x) <- value
```

```
## S3 replacement method for class 'ff'
vmode(x) <- value
  regtest.vmode()
```

### Arguments

x	any object
value	a vmode from .vmode
...	The ... don't have a function yet, they are only defined to keep the generic flexible.

### Details

vmode is generic with default and ff methods. The following meta data vectors can be queried by .vmode or .ffmode:

.vmode	virtual mode
.vunsigned	TRUE if unsigned vmode
.vvalues	number of possible values (incl. NA)
.vimplemented	TRUE if this vmode is available in ff (initialized .onLoad and stored in <a href="#">globalenv</a> )
.rammode	storage mode of this vmode
.ffmode	integer used to code the vmode in C-code
.vvalues	number of possible integers incl. NA in this vmode (or NA for other vmodes)
.vmin	min integer in this vmode (or NA for other vmodes)
.vmax	max integer in this vmode (or NA for other vmodes)
.vNA	NA or 0 if no NA for this vmode
.rambytes	bytes needed in ram
.ffbytes	bytes needed by ff on disk
.vcoerceable	list of vectors with those vmodes that can absorb this vmode

the following functions relate to vmode:

<a href="#">vector.vmode</a>	creating (ram) vector of some vmode
<a href="#">as.vmode</a>	generic for coercing to some vmode (dropping other attributes)
<a href="#">vmode&lt;-</a>	generic for coercing to some vmode (keeping other attributes)
<a href="#">maxffmode</a>	determine lowest .ffmode that can absorb all input vmodes without information loss

some of those call the vmode-specific functions:

creation	coercion	vmode description
<a href="#">boolean</a>	<a href="#">as.boolean</a>	1 bit logical without NA
<a href="#">logical</a>	<a href="#">as.logical</a>	2 bit logical with NA
<a href="#">quad</a>	<a href="#">as.quad</a>	2 bit unsigned integer without NA
<a href="#">nibble</a>	<a href="#">as.nibble</a>	4 bit unsigned integer without NA
<a href="#">byte</a>	<a href="#">as.byte</a>	8 bit signed integer with NA
<a href="#">ubyte</a>	<a href="#">as.ubyte</a>	8 bit unsigned integer without NA



short	as.short	16 bit signed integer with NA
ushort	as.ushort	16 bit unsigned integer without NA
integer	as.integer	32 bit signed integer with NA
single	as.single	32 bit float
double	as.double	64 bit float
complex	as.complex	2x64 bit float
raw	as.raw	8 bit unsigned char
character	as.character	character

**Value**

vmode returns a character scalar from .vmode or "NULL" for NULL  
 rambytes returns a vector of byte counts required by each of the vmodes

**Note**

regtest.vmode checks correctness of some vmode features

**Author(s)**

Jens Oehlschlägel

**See Also**

[ff](#), [storage.mode](#), [mode](#)

**Examples**

```
data.frame(.vmode=.vmode, .vimplemented=.vimplemented, .rammode=.rammode, .ffmode=.ffmode
, .vmin=.vmin, .vmax=.vmax, .vNA=.vNA, .rambytes=.rambytes, .ffbytes=.ffbytes)
vmode(1)
vmode(1L)
.vcoerceable[["byte"]]
.vcoerceable[["ubyte"]]
```

---

vmode.ffdf

*Virtual storage mode of ffdf*


---

**Description**

Function vmode returns the virtual storage mode of each ffdf column

**Usage**

```
## S3 method for class 'ffdf'
vmode(x, ...)
```

**Arguments**

x            [ffdf](#)  
 ...            ignored

**Value**

a character vector with one element for each column

**Author(s)**

Jens Oehlschlägel

**See Also**

[vmode](#), [ffdf](#)

**Examples**

```
vmode(as.ffdf(data.frame(a=as.double(1:26), b=letters)))
gc()
```

---

 vt

*Virtual transpose*


---

**Description**

The vt generic does a matrix or array transpose by modifying [virtual](#) attributes rather than by physically copying matrix elements.

**Usage**

```
vt(x, ...)
## S3 method for class 'ff'
vt(x, ...)
## Default S3 method:
vt(x, ...)
## S3 method for class 'ff'
t(x)
```

**Arguments**

x            an ff or ram object  
 ...            further arguments (not used)

## Details

The `vt.ff` method does transpose through reversing `dim.ff` and `dimorder`. The `vt.default` method is a wrapper to the standard transpose `t`.

The `t.ff` method creates a transposed `clone`.

If `x` has a virtual window `vw` defined, `vt.ff` returns an `ff` object with a transposed virtual window, the `t.ff` method return a transposed clone of the virtual window content only.

## Value

an object that behaves like a transposed matrix

## Author(s)

Jens Oehlschlägel

## See Also

`dim.ff`, `vw`, `virtual`

## Examples

```
x <- ff(1:20, dim=c(4,5))
x
vt(x)
y <- t(x)
y
vw(x) <- cbind(c(1,3,0),c(1,4,0))
x
vt(x)
y <- t(x)
y
rm(x,y); gc()
```

## Description

The virtual window `vw` function allows one to define a virtual window into an `ff_vector` or `ff_array`. The `ff` object will behave like a smaller array and it is mapped into the specified region of the complete array. This allows for example to execute recursive divide and conquer algorithms that work on parts of the full object, without the need to repeatedly create subfiles.

**Usage**

```

vw(x, ...)
vw(x, ...) <- value
## S3 method for class 'ff'
vw(x, ...)
## Default S3 method:
vw(x, ...)
## S3 replacement method for class 'ff_vector'
vw(x, ...) <- value
## S3 replacement method for class 'ff_array'
vw(x, ...) <- value

```

**Arguments**

x	an ff_vector or ff_array
...	further arguments (not used)
value	a vector or matrix with an Offset, Window and Rest component, see details and examples

**Details**

Each dimension of an ff array (or vector) is decomposed into three components, an invisible Offset, a visible Window and an invisible Rest. For each dimension the sum of the vw components must match the dimension (or length). For an ff\_vector, vw is simply a vector[1:3], for an array it is a matrix[1:3,1:length(dim(x))]. vw is a [virtual](#) attribute.

**Value**

NULL or a vw specification, see details

**Author(s)**

Jens Oehlschlägel

**See Also**

[length.ff](#), [dim.ff](#), [virtual](#)

**Examples**

```

x <- ff(1:26, names=letters)
y <- x
vw(x) <- c(0, 13, 13)
vw(y) <- c(13, 13, 0)
x
y
x[1] <- -1
y[1] <- -2

```

```

vw(x) <- NULL
x[]

z <- ff(1:24, dim=c(4,6), dimnames=list(letters[1:4], LETTERS[1:6]))
z
vw(z) <- rbind(c(1,1), c(2,4), c(1,1))
z

rm(x,y,z); gc()

```

---

write.table.ffdf

*Exporting csv files from ff data.frames*


---

## Description

Function `write.table.ffdf` writes a `ffdf` object to a separated flat file, very much like (and using) `write.table`. It can also work with any convenience wrappers like `write.csv` and provides its own convenience wrapper (e.g. `write.csv.ffdf`) for R's usual wrappers.

## Usage

```

write.table.ffdf(x = NULL
, file, append = FALSE
, nrows = -1, first.rows = NULL, next.rows = NULL
, FUN = "write.table", ...
, transFUN = NULL
, BATCHBYTES = getOption("ffbatchbytes")
, VERBOSE = FALSE
)
write.csv.ffdf(...)
write.csv2.ffdf(...)
write.csv(...)
write.csv2(...)

```

## Arguments

<code>x</code>	a <code>ffdf</code> object which to export to the separated file
<code>file</code>	either a character string naming a file or a connection open for writing. "" indicates output to the console.
<code>append</code>	logical. Only relevant if <code>file</code> is a character string. If <code>TRUE</code> , the output is appended to the file. If <code>FALSE</code> , any existing file of the name is destroyed.
<code>nrows</code>	integer: the maximum number of rows to write in (includes <code>first.rows</code> in case a 'first' chunk is read) Negative and other invalid values are ignored.
<code>first.rows</code>	the number of rows to write with the first chunk (default: <code>next.rows</code> )
<code>next.rows</code>	integer: number of rows to write in further chunks, see details. By default calculated as <code>BATCHBYTES %% sum(.rambytes[vmode(x)])</code>

FUN	character: name of a function that is called for writing each chunk, see <a href="#">write.table</a> , <a href="#">write.csv</a> , etc.
...	further arguments, passed to FUN in <code>write.table.ffdf</code> , or passed to <code>write.table.ffdf</code> in the convenience wrappers
transFUN	NULL or a function that is called on each data.frame chunk before writing with FUN (for filtering, transformations etc.)
BATCHBYTES	integer: bytes allowed for the size of the <code>data.frame</code> storing the result of reading one chunk. Default <code>getOption("ffbatchbytes")</code> .
VERBOSE	logical: TRUE to verbose timings for each processed chunk (default FALSE)

### Details

`write.table.ffdf` has been designed to export very large `ffdf` objects to separated flatfiles in chunks. The first chunk is potentially written with `col.names`. Further chunks are appended. `write.table.ffdf` has been designed to behave as much like [write.table](#) as possible. However, note the following differences:

1. by default `row.names` are only written if the `ffdf` has `row.names`.

### Value

`invisible`

### Note

`write.csv` and `write.csv2` have been fixed in order to suppress `col.names` if `append=TRUE` is passed. Note also that `write.table.ffdf` passes `col.names=FALSE` for all chunks following the first chunk - but not so for `FUN="write.csv"` and `FUN="write.csv2"`.

### Author(s)

Jens Oehlschlägel, Christophe Dutang

### See Also

[read.table.ffdf](#), [write.table](#), [ffdf](#)

### Examples

```
x <- data.frame(log=rep(c(FALSE, TRUE), length.out=26), int=1:26, dbl=1:26 + 0.1
, fac=factor(letters), ord=ordered(LETTERS), dct=Sys.time()+1:26
, dat=seq(as.Date("1910/1/1"), length.out=26, by=1))
ffx <- as.ffdf(x)

csvfile <- tempPathFile(path=getOption("fftempdir"), extension="csv")

write.csv.ffdf(ffx, file=csvfile)
write.csv.ffdf(ffx, file=csvfile, append=TRUE)

ffx <- read.csv.ffdf(file=csvfile, header=TRUE)
```

```

, colClasses=c(ord="ordered", dct="POSIXct", dat="Date"))

  rm(ffx, ffy); gc()
  unlink(csvfile)

## Not run:
# Attention, this takes very long
vmodes <- c(log="boolean", int="byte", dbl="single"
, fac="short", ord="short", dct="single", dat="single")

message("create a ffdf with 7 columns and 78 mio rows")
system.time({
  x <- data.frame(log=rep(c(FALSE, TRUE), length.out=26), int=1:26, dbl=1:26 + 0.1
, fac=factor(letters), ord=ordered(LETTERS), dct=Sys.time()+1:26
, dat=seq(as.Date("1910/1/1"), length.out=26, by=1))
  x <- do.call("rbind", rep(list(x), 10))
  x <- do.call("rbind", rep(list(x), 10))
  x <- do.call("rbind", rep(list(x), 10))
  x <- do.call("rbind", rep(list(x), 10))
  ffx <- as.ffdf(x, vmode = vmodes)
  for (i in 2:300){
    message(i, "\n")
    last <- nrow(ffx) + nrow(x)
    first <- last - nrow(x) + 1L
    nrow(ffx) <- last
    ffx[first:last,] <- x
  }
})

csvfile <- tempPathFile(path=getOption("fftempdir"), extension="csv")

write.csv.ffdf(ffx, file=csvfile, VERBOSE=TRUE)
ffx <- read.csv.ffdf(file=csvfile, header=TRUE
, colClasses=c(ord="ordered", dct="POSIXct", dat="Date")
, asffdf_args=list(vmode = vmodes), VERBOSE=TRUE)

rm(ffx, ffy); gc()
unlink(csvfile)

## End(Not run)

```

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