# Package 'bigmemory' 

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Description Create, store, access, and manipulate massive matrices.
Matrices are allocated to shared memory and may use memory-mapped files. Packages 'biganalytics', 'bigtabulate', 'synchronicity', and 'bigalgebra' provide advanced functionality.

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## $R$ topics documented:

bigmemory-package ..... 2
as.big.matrix-methods ..... 5
as.matrix,big.matrix-method ..... 5
big.matrix ..... 6
big.matrix-class ..... 10
deepcopy ..... 12
describe,big.matrix-method ..... 14
descriptor-class ..... 15
dim,big.matrix-method ..... 17
dimnames,big.matrix-method ..... 17
Extract,big.matrix ..... 18
flush ..... 20
GetMatrixSize ..... 21
head,big.matrix-method ..... 21
is.float ..... 22
is.float,numeric-method ..... 22
is.sub.big.matrix ..... 22
length,big.matrix-method ..... 24
morder ..... 24
mwhich ..... 26
mwhich-methods ..... 28
ncol,big.matrix-method ..... 28
print,big.matrix-method ..... 29
typeof,big.matrix-method ..... 29
write.big.matrix ..... 30
Index ..... 34

bigmemory-package Manage massive matrices with shared memory and memory-mapped
files.

## Description

Create, store, access, and manipulate massive matrices. Matrices are, by default, allocated to shared memory and may use memory-mapped files. Packages biganalytics, synchronicity, bigalgebra, and bigtabulate provide advanced functionality. Access to and manipulation of a big.matrix object is exposed in an S 4 class whose interface is similar to that of a matrix. Use of these packages in parallel environments can provide substantial speed and memory efficiencies. bigmemory also provides a C++ framework for the development of new tools that can work both with big.matrix and native matrix objects.

## Details

Index of functions/methods (grouped in a friendly way):

```
big.matrix, filebacked.big.matrix, as.big.matrix
    is.big.matrix, is.separated, is.filebacked
    describe, attach.big.matrix, attach.resource
    sub.big.matrix, is.sub.big.matrix
    dim, dimnames, nrow, ncol, print, head, tail, typeof, length
    read.big.matrix, write.big.matrix
    mwhich
    morder, mpermute
    deepcopy
    flush
```

Multi-gigabyte data sets challenge and frustrate users, even on well-equipped hardware. Use of C/C++ can provide efficiencies, but is cumbersome for interactive data analysis and lacks the flexibility and power of 's rich statistical programming environment. The package bigmemory and associated packages biganalytics, synchronicity, bigtabulate, and bigalgebra bridge this gap, implementing massive matrices and supporting their manipulation and exploration. The data structures may be allocated to shared memory, allowing separate processes on the same computer to share access to a single copy of the data set. The data structures may also be file-backed, allowing users to easily manage and analyze data sets larger than available RAM and share them across nodes of a cluster. These features of the Bigmemory Project open the door for powerful and memory-efficient parallel analyses and data mining of massive data sets.

This project (bigmemory and its sister packages) is still actively developed, although the design and current features can be viewed as "stable." Please feel free to email us with any questions: bigmemoryauthors@gmail.com.

## Memory considerations

For obvious reasons memory that the big.matrix uses is managed outside the R memory pool available to the garbage collector and the memory occupied by the big.matrix is not visible to the R. This has subtle implications:

- Memory usage is not visible via general $R$ functions (e.g. the $g c()$ function)
- Garbage collector is mislead by the very small memory footprint of the big.matrix object (which acts merely as a pointer to the external memory structure), which can result in much less eagerness to garbage-collect the unused big.memory objects. After removing a last reference to a big big. matrix, user should manually run gc() to reclaim the memory.
- Attaching the description of already finalized big.matrix and accessing this object will result in undefined behavior, which simply means it will crash the current R session with no hope of saving the data in it. To prevent R from de-allocating (finalizing) the matrices, user should keep at least one big.memory object somewhere in R memory in at least one R session on the current machine.
- Abruptly closed R (using e.g. task manager) will not have a chance to finalize the big.matrix objects, which will result in a memory leak, as the big.matrices will remain in the memory (perhaps under obfuscated names) with no easy way to reconnect R to them.


## Note

Various options are available. options(bigmemory.typecast.warning) can be set to avoid annoying warnings that might occur if, for example, you assign objects (typically type double) to char, short, or integer big.matrix objects. options(bigmemory.print.warning) protects against extracting and printing a massive matrix (which would involve the creation of a second massive copy of the matrix). options(bigmemory.allow.dimnames) by default prevents the setting of dimnames attributes, because they aren't allocated to shared memory and changes will not be visible across processes. options(bigmemory.default.type) is "double" be default (a change in default behavior as of 4.1.1) but may be changed by the user.
Note that you can't simply use a big. matrix with many (most) existing functions (e.g. lm, kmeans). One nice exception is split, because this function only accesses subsets of the matrix.

## Author(s)

Michael J. Kane, John W. Emerson, Peter Haverty, and Charles Determan Jr.
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## See Also

For example, big.matrix, mwhich, read.big.matrix

## Examples

```
# Our examples are all trivial in size, rather than burning huge amounts
# of memory.
x <- big.matrix(5, 2, type="integer", init=0,
        dimnames=list(NULL, c("alpha", "beta")))
x
x[1:2,]
x[,1] <- 1:5
x[,"alpha"]
colnames(x)
options(bigmemory.allow.dimnames=TRUE)
colnames(x) <- NULL
x[,]
```

as.big.matrix-methods Create a "big.matrix" from a matrix or vector.

## Description

Create a big.matrix from a matrix or vector or data. frame; a vector will result in a big. matrix with one column. A data frame will have character vectors converted to factors, and then all factors converted to numeric factor levels. All labels or character values will be lost.

## Methods

```
signature(x = "matrix") ...
signature(x = "vector") ...
signature(x = "data.frame") ...
```

```
as.matrix,big.matrix-method
        Convert to base R matrix
```


## Description

Extract values from a big.matrix object and convert to a base R matrix object

## Usage

\#\# S4 method for signature 'big.matrix'
as.matrix(x)

## Arguments

x
A big.matrix object
big.matrix The core "big.matrix" operations.

## Description

Create a big.matrix (or check to see if an object is a big.matrix, or create a big.matrix from a matrix, and so on). The big.matrix may be file-backed.

## Usage

```
big.matrix(
    nrow,
    ncol,
    type = options()$bigmemory.default.type,
    init = NULL,
    dimnames = NULL,
    separated = FALSE,
    backingfile = NULL,
    backingpath = NULL,
    descriptorfile = NULL,
    binarydescriptor = FALSE,
    shared = options()$bigmemory.default.shared
)
```

    filebacked.big.matrix(
        nrow,
        ncol,
        type = options()\$bigmemory.default.type,
        init = NULL,
        dimnames = NULL,
        separated = FALSE,
        backingfile = NULL,
        backingpath = NULL,
        descriptorfile \(=\) NULL,
        binarydescriptor = FALSE
    )
    as.big.matrix(
        x,
        type = NULL,
        separated = FALSE,
        backingfile = NULL,
        backingpath = NULL,
        descriptorfile = NULL,
        binarydescriptor = FALSE,
        shared \(=\) options()\$bigmemory.default.shared
    )
    ```
is.big.matrix(x)
## S4 method for signature 'big.matrix'
is.big.matrix(x)
## S4 method for signature 'ANY'
is.big.matrix(x)
is.separated(x)
## S4 method for signature 'big.matrix'
is.separated(x)
is.filebacked(x)
## S4 method for signature 'big.matrix'
is.filebacked(x)
shared.name(x)
## S4 method for signature 'big.matrix'
shared.name(x)
file.name(x)
## S4 method for signature 'big.matrix'
file.name(x)
dir.name(x)
## S4 method for signature 'big.matrix'
dir.name(x)
is.shared(x)
## S4 method for signature 'big.matrix'
is.shared(x)
is.readonly(x)
## S4 method for signature 'big.matrix'
is.readonly(x)
is.nil(address)
```


## Arguments

nrow number of rows.

| ncol | number of columns. <br> type type of the atomic element (options()\$bigmemory. default. type by de- <br> fault - "double" - but can be changed by the user to "integer", "short", or <br> "char"). |
| :--- | :--- |
| a scalar value for initializing the matrix (NULL by default to avoid unnecessary |  |
| time spent doing the initializing). |  |

## Details

A big.matrix consists of an object in $R$ that does nothing more than point to the data structure implemented in C++. The object acts much like a traditional R matrix, but helps protect the user from many inadvertent memory-consuming pitfalls of traditional R matrices and data frames.
There are two big.matrix types which manage data in different ways. A standard, shared big.matrix is constrained to available RAM, and may be shared across separate R processes. A file-backed big. matrix may exceed available RAM by using hard drive space, and may also be shared across processes. The atomic types of these matrices may be double, integer, short, or char (8, 4, 2, and 1 bytes, respectively).

If $x$ is a big.matrix, then $x[1: 5$,$] is returned as an R$ matrix containing the first five rows of $x$. If $x$ is of type double, then the result will be numeric; otherwise, the result will be an integer $R$ matrix. The expression $x$ alone will display information about the $R$ object (e.g. the external pointer) rather than evaluating the matrix itself (the user should try $x[$,$] with extreme caution,$ recognizing that a huge $R$ matrix will be created).

If $x$ has a huge number of rows and/or columns, then the use of rownames and/or colnames will be extremely memory-intensive and should be avoided. If $x$ has a huge number of columns and separated=TRUE is used (this isn't typically recommended), the user might want to store the transpose as there is overhead of a pointer for each column in the matrix. If separated is TRUE, then the memory is allocated into separate vectors for each column. Use this option with caution if you have a large number of columns, as shared-memory segments are limited by OS and hardware combinations. If separated is FALSE, the matrix is stored in traditional column-major format. The function is. separated() returns the separation type of the big.matrix.

When a big.matrix, $x$, is passed as an argument to a function, it is essentially providing call-byreference rather than call-by-value behavior. If the function modifies any of the values of $x$, the changes are not limited in scope to a local copy within the function. This introduces the possibility of side-effects, in contrast to standard R behavior.

A file-backed big.matrix may exceed available RAM in size by using a file cache (or possibly multiple file caches, if separated=TRUE). This can incur a substantial performance penalty for such large matrices, but less of a penalty than most other approaches for handling such large objects. A side-effect of creating a file-backed object is not only the file-backing(s), but a descriptor file (in the same directory) that is needed for subsequent attachments (see attach.big.matrix).

Note that we do not allow setting or changing the dimnames attributes by default; such changes would not be reflected in the descriptor objects or in shared memory. To override this, set options(bigmemory. allow. dimna

It should also be noted that a user can create an "anonymous" file-backed big.matrix by specifying "" as the filebacking argument. In this case, the backing resides in the temporary directory and a descriptor file is not created. These should be used with caution since even anonymous backings use disk space which could eventually fill the hard drive. Anonymous backings are removed either manually, by a user, or automatically, when the operating system deems it appropriate.

Finally, note that as.big.matrix can coerce data frames. It does this by making any character columns into factors, and then making all factors numeric before forming the big. matrix. Level labels are not preserved and must be managed by the user if desired.

## Value

A big.matrix is returned (for big.matrix and filebacked.big.matrix, and as.big.matrix), and TRUE or FALSE for is.big.matrix and the other functions.

## Author(s)

John W. Emerson and Michael J. Kane [bigmemoryauthors@gmail.com](mailto:bigmemoryauthors@gmail.com)

## References

The Bigmemory Project: http://www.bigmemory.org/.

## See Also

bigmemory, and perhaps the class documentation of big.matrix; attach.big.matrix and describe. Sister packages biganalytics, bigtabulate, synchronicity, and bigalgebra provide advanced functionality.

## Examples

```
x <- big.matrix(10, 2, type='integer', init=-5)
options(bigmemory.allow.dimnames=TRUE)
colnames(x) <- c("alpha", "beta")
is.big.matrix(x)
dim(x)
colnames(x)
rownames(x)
x[,]
x[1:8,1] <- 11:18
colnames(x) <- NULL
x[,]
# The following shared memory example is quite silly, as you wouldn't
# likely do this in a single R session. But if zdescription were
# passed to another R session via SNOW, foreach, or even by a
# simple file read/write, then the attach.big.matrix() within the
# second R process would give access to the same object in memory.
# Please see the package vignette for real examples.
z <- big.matrix(3, 3, type='integer', init=3)
z[,]
dim(z)
z[1,1] <- 2
z[,]
zdescription <- describe(z)
zdescription
y <- attach.big.matrix(zdescription)
y[,]
y
z
y[1,1]<--100
y[,]
z[,]
```

big.matrix-class Class "big.matrix"

## Description

The big.matrix class is designed for matrices with elements of type double, integer, short, or char. A big.matrix acts much like a traditional $R$ matrix, but helps protect the user from many inadvertent memory-consuming pitfalls of traditional $R$ matrices and data frames. The objects are allocated to shared memory, and if file-backing is used they may exceed virtual memory in size. Sadly, 32-bit operating system constraints - largely Windows and some MacOS versions -will be a limiting factor with file-backed matrices; 64-bit operating systems are recommended.

## Objects from the Class

Unlike many R objects, objects should not be created by calls of the form new("big.matrix", ...). The functions big.matrix() and filebacked.big.matrix() are intended for the user.

## Slots

address: Object of class "externalptr" points to the memory location of the $\mathrm{C}++$ data structure.

## Methods

As you would expect:

```
    signature(x = "big.matrix", i = "ANY", j = "ANY"): ...
[K<-signature(x = "big.matrix", i = "ANY", j = "missing"): ...
    [<- signature(x = "big.matrix", i = "missing", j = "ANY"): ...
    [<- signature(x = "big.matrix", i = "missing", j = "missing"): ...
    [<- signature(x = "big.matrix", i = "matrix", j = "missing"): ...
    [ signature(x = "big.matrix", i = "ANY", j = "ANY", drop = "missing"): ...
    [ signature(x = "big.matrix", i = "ANY", j = "ANY", drop = "logical"): ...
    [ signature(x = "big.matrix", i = "ANY", j = "missing", drop = "missing"): ...
    [ signature(x = "big.matrix", i = "ANY", j = "missing", drop = "logical"): ...
    [ signature(x = "big.matrix", i = "matrix", j = "missing", drop = "logical"): ...
    [ signature(x = "big.matrix", i = "missing", j = "ANY", drop = "missing"): ...
    [ signature(x = "big.matrix", i = "missing", j = "ANY", drop = "logical"): ...
    [ signature(x = "big.matrix", i = "missing", j = "missing", drop = "missing"): ...
    [ signature(x = "big.matrix", i = "missing", j = "missing", drop = "logical"): ...
```

        The following are probably more interesting:
    describe signature ( \(\mathrm{x}=\) "big.matrix"): provide necessary and sufficient information for the
        sharing or re-attaching of the object.
    dim signature ( \(x=\) "big.matrix"): returns the dimension of the big.matrix.
    length signature( \(x=\) "big.matrix"): returns the product of the dimensions of the big.matrix.
    dimnames<- signature( \(x=\) "big.matrix", value = "list"): set the row and column names,
        prohibited by default (see bigmemory to override).
    dimnames signature ( $x=$ "big.matrix"): get the row and column names.
head signature ( $x=$ "big.matrix"): get the first 6 (or $n$ ) rows.
as.matrix signature ( $x=$ "big.matrix"): coerce abig.matrix to a matrix.
is.big.matrix signature ( $x=$ "big.matrix"): return TRUE if it's a big.matrix.
is.filebacked signature ( $x=$ "big.matrix"): return TRUE if there is a file-backing.
is.separated signature ( $x=$ "big.matrix") : return TRUE if the big.matrix is organized as a separated column vectors.
is.sub.big.matrix signature ( $x=$ "big.matrix"): return TRUE if this is a sub-matrix of a big.matrix.
ncol signature( $x=$ "big.matrix"): returns the number of columns.
nrow signature ( $x=$ "big.matrix"): returns the number of rows.
print signature( $x=$ "big.matrix"): a traditional print() is intentionally disabled, and returns head $(x)$ unless options()\$bm. print.warning==FALSE; in this case, print( $x[$,$] ) is the$ result, which could be very big!
sub.big.matrix signature( $x=$ "big.matrix"): for contiguous submatrices.
tail signature ( $x=$ "big.matrix") : returns the last 6 (or $n$ ) rows.
typeof signature( $x=$ "big.matrix"): return the type of the atomic elements of the big.matrix.
write.big.matrix signature(bigMat = "big.matrix", fileName = "character"): produce an ASCII file from the big.matrix.
apply signature ( $x=$ "big.matrix") : apply () where MARGIN may only be 1 or 2 , but otherwise conforming to what you would expect from apply().

## Author(s)

Michael J. Kane and John W. Emerson [bigmemoryauthors@gmail.com](mailto:bigmemoryauthors@gmail.com)

## See Also

big.matrix

## Examples

showClass("big.matrix")
deepcopy Produces a physical copy of a "big.matrix"

## Description

This is needed to make a duplicate of a big. matrix, with the new copy optionally filebacked.

```
Usage
    deepcopy(
        x,
        cols = NULL,
        rows = NULL,
    y = NULL,
    type = NULL,
    separated = NULL,
    backingfile = NULL,
    backingpath = NULL,
    descriptorfile = NULL,
    binarydescriptor = FALSE,
    shared = options()$bigmemory.default.shared
    )
```


## Arguments

X
cols
rows possible subset of rows for the deepcopy; could be numeric, named, or logical.
$y \quad$ optional destination object (matrix or big.matrix); if not specified, abig.matrix will be created.
type preferably specified, "integer" for example.
separated use separated column organization of the data instead of column-major organization; use with caution if the number of columns is large.
backingfile the root name for the file(s) for the cache of $x$.
backingpath the path to the directory containing the file-backing cache.
descriptorfile we recommend specifying this for file-backing.
binarydescriptor
the flag to specify if the binary RDS format should be used for the backingfile description, for subsequent use with attach.big.matrix; if NULL of FALSE, the dput () file format is used.
shared TRUE by default, and always TRUE if the big.matrix is file-backed. For a nonfilebacked big.matrix, shared=FALSE uses non-shared memory, which can be more stable for large (say, $>50 \backslash$ fail in such cases due to exhausted sharedmemory resources in the system.

## Details

This is needed to make a duplicate of a big.matrix, because traditional syntax would only copy the object (the pointer to the big.matrix rather than the big.matrix itself). It can also make a copy of only a subset of columns.

## Value

abig.matrix.

## See Also

```
    big.matrix
```


## Examples

```
x <- as.big.matrix(matrix(1:30, 10, 3))
y <- deepcopy(x, -1) # Don't include the first column.
x
y
head(x)
head(y)
```

```
describe,big.matrix-method
```

                                    The basic "big.matrix" operations for sharing and re-attaching.
    
## Description

The describe function returns the information needed by attach.big.matrix to reference a shared or file-backed big.matrix object. The attach.big.matrix and attach. resource functions create a new big.matrix object based on the descriptor information referencing previously allocated shared-memory or file-backed matrices.

## Usage

```
## S4 method for signature 'big.matrix'
describe(x)
    attach.big.matrix(obj, ...)
```


## Arguments

x
obj an object as returned by describe() or, optionally, the filename of the descriptor for a filebacked matrix, assumed to be in the directory specified by the path (if one is provided)
... possibly path which gives the path where the descriptor and/or filebacking can be found

## Details

The describe function returns a list of the information needed to attach to a big.matrix object. A descriptor file is automatically created when a new filebacked big.matrix is created.

## Value

describe returns a list of of the information needed to attach to a big. matrix object.
attach.big.matrix return a new instance of type big.matrix corresponding to a shared-memory or file-backed big. matrix.

## Author(s)

Michael J. Kane and John W. Emerson <bigmemoryauthors@gmail. com>

## See Also

bigmemory, big.matrix, or the class documentation big.matrix.

## Examples

```
    # The example is quite silly, as you wouldn't likely do this in a
    # single R session. But if zdescription were passed to another R session
    # via SNOW, foreach, or even by a simple file read/write,
    # then the attach of the second R process would give access to the
    # same object in memory. Please see the package vignette for real examples.
    z <- big.matrix(3, 3, type='integer', init=3)
    z[,]
    dim(z)
    z[1,1] <- 2
    z[,]
    zdescription <- describe(z)
    zdescription
    y <- attach.big.matrix(zdescription)
    y[,]
    y
    z
    zz <- attach.resource(zdescription)
    zz[1,1] <- -100
    y[,]
    z[,]
```

    descriptor-class Class "big.matrix.descriptor"
    
## Description

An object of this class contains necessary and sufficient information to "attach" a shared or filebacked big.matrix.

## Usage

```
## S4 method for signature 'big.matrix.descriptor'
sub.big.matrix(
    x,
    firstRow = 1,
    lastRow = NULL,
    firstCol = 1,
    lastCol = NULL,
    backingpath = NULL
)
## S4 method for signature 'character'
attach.resource(obj, ...)
    ## S4 method for signature 'big.matrix.descriptor'
    attach.resource(obj, ...)
```


## Arguments

| x | A descriptor object |
| :--- | :--- |
| firstRow | the first row of the submatrix |
| lastRow | the last row of the submatrix if not NULL <br> firstCol <br> lastCol <br> backingpath <br> obj |
| of the submatrix if not NULL <br> required path to the filebacked object, if applicable |  |
| $\ldots$ | The filename of the descriptor for a filebacked matrix, assumed to be in the <br> directory specified |
| possibly path which gives the path where the descriptor and/or filebacking can |  |
| be found. |  |

## Objects from the Class

Objects should not be created by calls of the form new("big.matrix.descriptor", ...), but should use the describe function.

## Slots

description: Object of class "list"; details omitted.

## Extends

Class "descriptor", directly.

## Methods

attach.resource signature(obj = "big.matrix.descriptor"): ...
sub.big.matrix signature(x = "big.matrix.descriptor"): ...

## Note

We provide attach. resource for convenience, but expect most users will prefer attach.big. matrix.

## Author(s)

John W. Emerson and Michael J. Kane

## References

Other types of descriptors are defined in package synchronicity.

See Also
See also attach.big.matrix.

## Examples

```
showClass("big.matrix.descriptor")
```

dim,big.matrix-method Dimensions of a big.matrix object

## Description

Retrieve the dimensions of a big.matrix object

## Usage

\#\# S4 method for signature 'big.matrix'
$\operatorname{dim}(x)$

## Arguments

x
A big.matrix object
dimnames, big.matrix-method
Dimnames of a big.matrix Object

## Description

Retrieve or set the dimnames of an object

## Usage

\#\# S4 method for signature 'big.matrix'
dimnames ( x )
\#\# S4 replacement method for signature 'big.matrix,list' dimnames $(x)$ <- value

## Arguments

X
value A possible value for dimnames ( $x$ )

Extract,big.matrix Extract or Replace

## Description

Extract or replace big.matrix elements

## Usage

\#\# S4 method for signature 'big.matrix,ANY,ANY,missing'
x[i, j, drop]
\#\# S4 method for signature 'big.matrix, ANY, ANY,logical'
x[i, j, drop]
\#\# S4 method for signature 'big.matrix,missing, ANY,missing'
x[i, j, drop]
\#\# S4 method for signature 'big.matrix,missing,ANY,logical'
x[i, j, drop]
\#\# S4 method for signature 'big.matrix,ANY,missing,missing'
x[i, j, ..., drop = TRUE]
\#\# S4 method for signature 'big.matrix,ANY,missing,logical'
$x[i, j, d r o p]$
\#\# S4 method for signature 'big.matrix,missing,missing,missing'
x[i, j, drop]
\#\# S4 method for signature 'big.matrix,missing,missing,logical'
x[i, j, drop]
\#\# S4 method for signature 'big.matrix,matrix,missing,missing'
x[i, j, drop]
\#\# S4 replacement method for signature 'big.matrix, numeric, numeric, ANY' $x[i, j]<-$ value
\#\# S4 replacement method for signature 'big.matrix, numeric,logical, ANY' $x[i, j]<-$ value
\#\# S4 replacement method for signature 'big.matrix,logical,numeric,ANY' x[i, j] <- value
\#\# S4 replacement method for signature 'big.matrix,logical,logical,ANY'
$x[i, j]$ <- value

```
## S4 replacement method for signature 'big.matrix,logical,character,ANY'
x[i, j] <- value
## S4 replacement method for signature 'big.matrix,numeric,character,ANY'
x[i, j] <- value
## S4 replacement method for signature 'big.matrix,missing,missing,ANY'
x[i, j] <- value
## S4 replacement method for signature 'big.matrix,missing,numeric,ANY'
x[i, j] <- value
## S4 replacement method for signature 'big.matrix,missing,logical,ANY'
x[i, j] <- value
## S4 replacement method for signature 'big.matrix,numeric,missing,numeric'
x[i, j, ...] <- value
## S4 replacement method for signature 'big.matrix,logical,missing,numeric'
x[i, j, ...] <- value
## S4 replacement method for signature 'big.matrix,numeric,missing,matrix'
x[i, j, ...] <- value
## S4 replacement method for signature 'big.matrix,logical,missing,matrix'
x[i, j, ...] <- value
## S4 replacement method for signature 'big.matrix,character,character,ANY'
x[i, j] <- value
## S4 replacement method for signature 'big.matrix,missing,character,ANY'
x[j] <- value
## S4 replacement method for signature 'big.matrix,character,missing,ANY'
x[i] <- value
## S4 replacement method for signature 'big.matrix,missing,missing,numeric'
x[i, j] <- value
## S4 replacement method for signature 'big.matrix,matrix,missing,numeric'
x[i, j] <- value
```


## Arguments

X
i
j

A big.matrix object
Indices specifying the rows
Indices specifying the columns

| drop | Logical indication if reduce to minimum dimensions |
| :--- | :--- |
| $\ldots$ | Additional arguments |
| value | typically an array-like R object of similar class |

flush Updating a big.matrix filebacking.

## Description

For a file-backed big.matrix object, flush() forces any modified information to be written to the file-backing.

## Usage

flush(con)
\#\# S4 method for signature 'big.matrix'
flush(con)

## Arguments

con filebacked big.matrix.

## Details

This function flushes any modified data (in RAM) of a file-backed big.matrix to disk. This may be useful for improving performance in cases where allowing the operating system to decide on flushing creates a bottleneck (likely near the threshold of available RAM).

## Value

TRUE or FALSE (invisible), indicating whether or not the flush was successful.

## Author(s)

John W. Emerson and Michael J. Kane

## Examples

```
temp_dir = tempdir()
if (!dir.exists(temp_dir)) dir.create(temp_dir)
x <- big.matrix(nrow=3, ncol=3, backingfile='flushtest.bin',
    descriptorfile='flushtest.desc', backingpath=temp_dir,
    type='integer')
x[1,1] <- 0
flush(x)
```

```
    GetMatrixSize big.matrix size
```


## Description

Returns the size of the created matrix in bytes

## Usage

GetMatrixSize(bigMat)

## Arguments

```
    bigMat abig.matrix object
```

    head, big.matrix-method
    Return First or Last Part of a big.matrix Object
    
## Description

Returns the first or last parts of a big. matrix object.

## Usage

```
## S4 method for signature 'big.matrix'
head(x, n = 6)
## S4 method for signature 'big.matrix'
tail(x, n = 6)
```


## Arguments

x
n

A big.matrix object
A single integer for the number of rows to return
is.float Check if Float

## Description

Check to see if the elements of a big.matrix object are floats.

## Usage

is.float(x)

## Arguments

X
An object to be evaluated if float

```
is.float,numeric-method
    Is Float?
```


## Description

Check if R numeric value has float flag

## Usage

\#\# S4 method for signature 'numeric'
is.float(x)

## Arguments

x
A numeric value

```
is.sub.big.matrix Submatrix support
```


## Description

This doesn't create a copy, it just provides a new version of the class which provides behavior for a contiguous submatrix of the big.matrix. Non-contiguous submatrices are not supported.

## Usage

```
is.sub.big.matrix(x)
## S4 method for signature 'big.matrix'
is.sub.big.matrix(x)
    sub.big.matrix(
        x,
        firstRow = 1,
        lastRow = NULL,
        firstCol = 1,
        lastCol = NULL,
        backingpath = NULL
)
## S4 method for signature 'big.matrix'
sub.big.matrix(
        x,
        firstRow = 1,
        lastRow = NULL,
        firstCol = 1,
        lastCol = NULL,
        backingpath = NULL
)
```


## Arguments

x
firstRow the first row of the submatrix.
lastRow the last row of the submatrix if not NULL.
firstCol the first column of the submatrix.
lastCol the last column of the submatrix if not NULL.
backingpath required path to the filebacked object, if applicable.

## Details

The sub.big.matrix function allows a user to create a big.matrix object that references a contiguous set of columns and rows of another big. matrix object.

The is.sub.big.matrix function returns TRUE if the specified argument is a sub.big.matrix object and return FALSE otherwise.

## Value

A big.matrix which is actually a submatrix of a larger big.matrix. It is not a physical copy. Only contiguous blocks may form a submatrix.

## Author(s)

John W. Emerson and Michael J. Kane

## See Also

big.matrix

## Examples

```
x <- big.matrix(10, 5, init=0, type="double")
x[,] <- 1:50
y <- sub.big.matrix(x, 2, 9, 2, 3)
y[,]
y[1,1] <- -99
x[,]
rm(x)
```

    length, big.matrix-method
    Length of a big.matrix object
    
## Description

Get the length of a big.matrix object

## Usage

```
## S4 method for signature 'big.matrix'
    length(x)
```


## Arguments



A big.matrix object

morder | Ordering and Permuting functions for big.matrix', and matrix" |
| :--- |
| objects |

## Description

The morder function returns a permutation of row indices which can be used to rearrange an object according to the values in the specified columns (a multi-column ordering). The mpermute function actually reorders the rows of a big.matrix or matrix based on an order vector or a desired ordering on a set of columns.
morder

## Usage

morder (x, cols, na.last $=$ TRUE, decreasing $=$ FALSE)
morderCols(x, rows, na.last $=$ TRUE, decreasing $=$ FALSE)
mpermute (x, order $=$ NULL, cols $=$ NULL, allow.duplicates $=$ FALSE, ...)
mpermuteCols(x, order = NULL, rows = NULL, allow.duplicates = FALSE, ...)

## Arguments

x
A big.matrix or matrix object with numeric values.
cols The columns of $x$ to get the ordering for or reorder on
na. last for controlling the treatment of NAs. If TRUE, missing values in the data are put last; if FALSE, they are put first; if NA, they are removed.
decreasing logical. Should the sort order be increasing or decreasing?
rows $\quad$ The rows of $x$ to get the ordering for or reorder on
order A vector specifying the reordering of rows, i.e. the result of a call to order or morder.
allow.duplicates
ff TRUE, allows a row to be duplicated in the resulting big.matrix or matrix (i.e. in this case, order would not need to be a permutation of 1 : $\operatorname{nrow}(x)$ ).
... optional parameters to pass to morder when cols is specified instead of just using order.

## Details

The morder function behaves similar to order, returning a permutation of 1 : $\mathrm{nrow}(x)$ which rearranges objects according to the values in the specified columns. However, morder takes a big. matrix or an $R$ matrix (with numeric type) and a set of columns (cols) with which to determine the ordering; morder does not incur the same memory overhead required by order, and runs more quickly.
The mpermute function changes the row ordering of a big.matrix or matrix based on a vector order or an ordering based on a set of columns specified by cols. It should be noted that this function has side-effects, that is x is changed when this function is called.

## Value

morder returns an ordering vector. mpermute returns nothing but does change the contents of $x$. This type of a side-effect is generally frowned upon in R, but we "break" the rules here to avoid memory overhead and improve performance.

## Author(s)

Michael J. Kane [bigmemoryauthors@gmail.com](mailto:bigmemoryauthors@gmail.com)

## See Also

order

## Examples

```
    m = matrix(as.double(as.matrix(iris)), nrow=nrow(iris))
    morder(m, 1)
    order(m[,1])
    m[order(m[,1]), 2]
    mpermute(m, cols=1)
    m[,2]
```

mwhich Expanded "which"-like functionality.

## Description

Implements which-like functionality for a big.matrix, with additional options for efficient comparisons (executed in $\mathrm{C}++$ ); also works for regular numeric matrices without the memory overhead.

## Usage

mwhich(x, cols, vals, comps, op = "AND")

## Arguments

x
a big.matrix (or a numeric matrix; see below).
cols a vector of column indices or names.
vals a list (one component for each of cols) of vectors of length 1 or 2 ; length 1 is used to test equality (or inequality), while vectors of length 2 are used for checking values in the range (-Inf and Inf are allowed). If a scalar or vector of length 2 is provided instead of a list, it will be replicated length(cols) times.
comps a list of operators (one component for each of cols), including 'eq', 'neq', 'le', 'lt', 'ge' and 'gt'. If a single operator, it will be replicated length(cols) times.
op the comparison operator for combining the results of the individual tests, either 'AND' or 'OR'.

## Details

To improve performance and avoid the creation of massive temporary vectors in $R$ when doing comparisons, mwhich() efficiently executes column-by-column comparisons of values to the specified values or ranges, and then returns the row indices satisfying the comparison specified by the op operator. More advanced comparisons are then possible (and memory-efficient) in R by doing set operations (union and intersect, for example) on the results of multiple mwhich() calls.
Note that NA is a valid argument in conjunction with 'eq' or 'neq', replacing traditional is.na() calls. And both -Inf and Inf can be used for one-sided inequalities.
If mwhich() is used with a regular numeric R matrix, we access the data directly and thus incur no memory overhead. Interested developers might want to look at our code for this case, which uses a handy pointer trick (accessor) in C++.

## Value

a vector of row indices satisfying the criteria.

## Author(s)

John W. Emerson [bigmemoryauthors@gmail.com](mailto:bigmemoryauthors@gmail.com)

## See Also

big.matrix, which

## Examples

```
x <- as.big.matrix(matrix(1:30, 10, 3))
options(bigmemory.allow.dimnames=TRUE)
colnames(x) <- c("A", "B", "C")
x[,]
x[mwhich(x, 1:2, list(c(2,3), c(11,17)),
        list(c('ge','le'), c('gt', 'lt')), 'OR'),]
x[mwhich(x, c("A","B"), list(c(2,3), c(11,17)),
        list(c('ge','le'), c('gt', 'lt')), 'AND'),]
# These should produce the same answer with a regular matrix:
y <- matrix(1:30, 10, 3)
y[mwhich(y, 1:2, list(c(2,3), c(11,17)),
    list(c('ge','le'), c('gt', 'lt')), 'OR'),]
y[mwhich(y, -3, list(c(2,3), c(11,17)),
    list(c('ge','le'), c('gt', 'lt')), 'AND'),]
x[1,1]<-NA
mwhich(x, 1:2, NA, 'eq', 'OR')
mwhich(x, 1:2, NA, 'neq', 'AND')
# Column 1 equal to 4 and/or column 2 less than or equal to 16:
mwhich(x, 1:2, list(4, 16), list('eq', 'le'), 'OR')
mwhich(x, 1:2, list(4, 16), list('eq', 'le'), 'AND')
# Column 2 less than or equal to 15:
mwhich(x, 2, 15, 'le')
# No NAs in either column, and column 2 strictly less than 15:
mwhich(x, c(1:2,2), list(NA, NA, 15), list('neq', 'neq', 'lt'), 'AND')
x <- big.matrix(4, 2, init=1, type="double")
x[1,1]<- Inf
mwhich(x, 1, Inf, 'eq')
mwhich(x, 1, 1, 'gt')
mwhich(x, 1, 1, 'le')
```

```
mwhich-methods Expanded "which"-like functionality.
```


## Description

Implements which-like functionality for a big.matrix, with additional options for efficient comparisons (executed in $\mathrm{C}++$ ); also works for regular numeric matrices without the memory overhead. test

## Methods

```
signature(x = "big.matrix=', cols = "ANY", vals = "ANY',", " comps = "ANY", op = 'character')
signature(x = 'big.matrix", cols = "ANY', vals = "ANY',", ' comps = "ANY', op = 'missing')
    ...
    signature(x = 'matrix', cols = "ANY", vals = "ANY",", ' comps = 'ANY', op = 'character'')
    ...
    signature(x = ''matrix', cols = "ANY', vals = "ANY'',', ' comps = 'ANY', op = ''missing'')
```

See Also
big.matrix, which, mwhich
ncol, big.matrix-method
The Number of Rows/Columns of a big.matrix

## Description

nrow and ncol return the number of rows or columns present in a big.matrix object.

## Usage

```
    ## S4 method for signature 'big.matrix'
```

    ncol ( \(x\) )
    \#\# S4 method for signature 'big.matrix'
    nrow (x)
    
## Arguments

x
A big.matrix object

## Value

An integer of length 1

```
print,big.matrix-method
```

Print Values

## Description

print will print out the elements within a big.matrix object.

## Usage

\#\# S4 method for signature 'big.matrix' print (x)

## Arguments

$x \quad$ A big.matrix object

## Note

By default, this will only return the head of a big.matrix to prevent console overflow. If you turn off the bigmemory.print.warning option then it will convert to a base R matrix and print all elements.

```
typeof,big.matrix-method
    The Type of a big.matrix Object
```


## Description

typeof returns the storage type of a big.matrix object

## Usage

\#\# S4 method for signature 'big.matrix'
typeof(x)

## Arguments

```
write.big.matrix File interface for a "big.matrix"
```


## Description

Create a big.matrix by reading from a suitably-formatted ASCII file, or write the contents of a big.matrix to a file.

## Usage

```
write.big.matrix(x, filename, row.names = FALSE, col.names = FALSE, sep = ",")
## S4 method for signature 'big.matrix,character'
write.big.matrix(x, filename, row.names = FALSE, col.names = FALSE, sep = ",")
read.big.matrix(
        filename,
        sep = ",",
        header = FALSE,
        col.names = NULL,
        row.names = NULL,
        has.row.names = FALSE,
        ignore.row.names = FALSE,
        type = NA,
        skip = 0,
        separated = FALSE,
        backingfile = NULL,
        backingpath = NULL,
        descriptorfile = NULL,
        binarydescriptor = FALSE,
        extraCols = NULL,
        shared = options()$bigmemory.default.shared
    )
    ## S4 method for signature 'character'
    read.big.matrix(
    filename,
    sep = ",",
    header = FALSE,
    col.names = NULL,
    row.names = NULL,
    has.row.names = FALSE,
    ignore.row.names = FALSE,
    type = NA,
    skip = 0,
    separated = FALSE,
    backingfile = NULL,
```

```
    backingpath = NULL,
    descriptorfile = NULL,
    binarydescriptor = FALSE,
    extraCols = NULL,
    shared = options()$bigmemory.default.shared
)
```


## Arguments

X
filename
row.names a vector of names, use them even if row names appear to exist in the file.
col.names a vector of names, use them even if column names exist in the file.
sep a field delimiter.
header if TRUE, the first line (after a possible skip) should contain column names.
has. row. names if TRUE, then the first column contains row names.
ignore. row.names
if TRUE when has.row. names==TRUE, the row names will be ignored.
type preferably specified, "integer" for example.
skip number of lines to skip at the head of the file.
separated use separated column organization of the data instead of column-major organization.
backingfile the root name for the file(s) for the cache of $x$.
backingpath the path to the directory containing the file backing cache.
descriptorfile the file to be used for the description of the filebacked matrix.
binarydescriptor
the flag to specify if the binary RDS format should be used for the backingfile description, for subsequent use with attach.big. matrix; if NULL of FALSE, the dput () file format is used.
extraCols the optional number of extra columns to be appended to the matrix for future use.
shared if TRUE, the resulting big.matrix can be shared across processes.

## Details

Files must contain only one atomic type (all integer, for example). You, the user, should know whether your file has row and/or column names, and various combinations of options should be helpful in obtaining the desired behavior.
When reading from a file, if type is not specified we try to make a reasonable guess for you without making any guarantees at this point. Unless you have really large integer values, we recommend you consider "short". If you have something that is essentially categorical, you might even be able use "char", with huge memory savings for large data sets.

Any non-numeric entry will be ignored and replaced with NA, so reading something that traditionally would be a data. frame won't cause an error. A warning is issued.

Wishlist: we'd like to provide an option to ignore specified columns while doing reads. Or perhaps to specify columns targeted for factor or character conversion to numeric values. Would you use such features? Email us and let us know!

## Value

a big.matrix object is returned by read.big.matrix, while write.big.matrix creates an output file (a path could be part of filename).

## Author(s)

John W. Emerson and Michael J. Kane <bigmemoryauthors@gmail. com>

## See Also

big.matrix

## Examples

```
# Without specifying the type, this big.matrix x will hold integers.
x <- as.big.matrix(matrix(1:10, 5, 2))
x[2,2] <- NA
x[,]
temp_dir = tempdir()
if (!dir.exists(temp_dir)) dir.create(temp_dir)
write.big.matrix(x, file.path(temp_dir, "foo.txt"))
# Just for fun, I'll read it back in as character (1-byte integers):
y <- read.big.matrix(file.path(temp_dir, "foo.txt"), type="char")
y[,]
# Other examples:
w <- as.big.matrix(matrix(1:10, 5, 2), type='double')
w[1,2] <- NA
w[2,2] <- -Inf
w[3,2] <- Inf
w[4,2] <- NaN
w[,]
write.big.matrix(w, file.path(temp_dir, "bar.txt"))
w <- read.big.matrix(file.path(temp_dir, "bar.txt"), type="double")
w[,]
w <- read.big.matrix(file.path(temp_dir, "bar.txt"), type="short")
w[,]
# Another example using row names (which we don't like).
x <- as.big.matrix(as.matrix(iris), type='double')
rownames(x) <- as.character(1:nrow(x))
head(x)
write.big.matrix(x, file.path(temp_dir, 'IrisData.txt'), col.names=TRUE,
                                    row.names=TRUE)
y <- read.big.matrix(file.path(temp_dir, "IrisData.txt"), header=TRUE,
```

```
    has.row.names=TRUE)
head(y)
# The following would fail with a dimension mismatch:
if (FALSE) y <- read.big.matrix(file.path(temp_dir, "IrisData.txt"),
                header=TRUE)
```


## Index

* classes
big.matrix, 6
big.matrix-class, 10
describe,big.matrix-method, 14
descriptor-class, 15
* methods
as.big.matrix-methods, 5
big.matrix, 6
deepcopy, 12
describe,big.matrix-method, 14
flush, 20
is.sub.big.matrix, 22
mwhich, 26
mwhich-methods, 28
write.big.matrix, 30
* package
bigmemory-package, 2
[,big.matrix, ANY, ANY, logical-method (Extract,big.matrix), 18
[, big.matrix, ANY, ANY, missing-method (Extract,big.matrix), 18
[,big.matrix, ANY, missing,logical-method (Extract,big.matrix), 18
[,big.matrix, ANY,missing,missing-method (Extract,big.matrix), 18
[,big.matrix,matrix,missing,missing-method (Extract,big.matrix), 18
[,big.matrix,missing, ANY, logical-method (Extract,big.matrix), 18
[,big.matrix,missing, ANY, missing-method (Extract,big.matrix), 18
[,big.matrix,missing,missing,logical-method (Extract,big.matrix), 18
[,big.matrix,missing,missing, missing-method (Extract,big.matrix), 18
[<-, big.matrix, ANY, ANY, ANY-method (Extract,big.matrix), 18
[<-, big.matrix, character, missing, ANY-method
(Extract,big.matrix), 18
[<-,big.matrix,logical, character, ANY-method (Extract,big.matrix), 18
[<-,big.matrix,logical,logical, ANY-method (Extract,big.matrix), 18
[<-,big.matrix,logical,missing, matrix-method (Extract,big.matrix), 18
[<-, big.matrix,logical, missing, numeric-method (Extract, big.matrix), 18
[<-, big.matrix, logical, numeric, ANY-method (Extract,big.matrix), 18
[<-, big.matrix, matrix, missing, numeric-method (Extract,big.matrix), 18
[<-,big.matrix,missing, character, ANY-method (Extract,big.matrix), 18
[<-, big.matrix,missing,logical, ANY-method (Extract,big.matrix), 18
[<-, big.matrix, missing, missing, ANY-method (Extract, big.matrix), 18
[<-, big.matrix,missing,missing, numeric-method (Extract,big.matrix), 18
[<-, big.matrix, missing, numeric, ANY-method (Extract,big.matrix), 18
[<-, big.matrix, numeric, character, ANY-method (Extract,big.matrix), 18
[<-,big.matrix, numeric, logical, ANY-method (Extract,big.matrix), 18
[<-, big.matrix, numeric, missing, matrix-method (Extract,big.matrix), 18
[<-, big.matrix, numeric, missing, numeric-method (Extract, big.matrix), 18
[<-, big.matrix, numeric, numeric, ANY-method (Extract,big.matrix), 18
as.big.matrix (big.matrix), 6
as.big.matrix, data.frame-method
(as.big.matrix-methods), 5
[<-, big.matrix, character, character, ANY-methodas.big.matrix, matrix-method (Extract,big.matrix), 18
(as.big.matrix-methods), 5
as.big.matrix, vector-method
(as.big.matrix-methods), 5
as.big.matrix-methods, 5
as.matrix,big.matrix-method, 5
attach.big.matrix, 8, 9, 13, 16, 31
attach.big.matrix
(describe, big.matrix-method),
14
attach.resource
(describe,big.matrix-method),
14
attach.resource,big.matrix.descriptor-method (descriptor-class), 15
attach.resource, character-method (descriptor-class), 15
big.matrix, 2, 4, 5, 6, 9, 12-15, 20, 23, 24,
26-28, 30-32
big.matrix-class, 10
big.matrix.descriptor-class
(descriptor-class), 15
bigmemory, 9, 11, 14
bigmemory (bigmemory-package), 2
bigmemory-package, 2
data.frame, 5
deepcopy, 12
describe, 9, 16
describe (describe, big.matrix-method),
14
describe,big.matrix-method, 14
descriptor, 16
descriptor-class, 15
dim, big.matrix-method, 17
dimnames, big.matrix-method, 17
dimnames<-, big.matrix,list-method
(dimnames, big.matrix-method), 17
dir.name (big.matrix), 6
dir.name, big.matrix-method
(big.matrix), 6
Extract, big.matrix, 18
file. name (big.matrix), 6
file.name,big.matrix-method
(big.matrix), 6
filebacked.big.matrix (big.matrix), 6
flush, 20
flush,big.matrix-method (flush), 20
GetMatrixSize, 21
head, big.matrix-method, 21
intersect, 26
is.big.matrix (big.matrix), 6
is.big.matrix, ANY-method (big.matrix), 6
is.big.matrix,big.matrix-method
(big.matrix), 6
is.filebacked (big.matrix), 6
is.filebacked,big.matrix-method
(big.matrix), 6
is.float, 22
is.float, numeric-method, 22
is.nil (big.matrix), 6
is.readonly (big.matrix), 6
is.readonly,big.matrix-method (big.matrix), 6
is.separated (big.matrix), 6
is.separated,big.matrix-method (big.matrix), 6
is.shared (big.matrix), 6
is.shared,big.matrix-method (big.matrix), 6
is.sub.big.matrix, 22
is.sub.big.matrix, big.matrix-method (is.sub.big.matrix), 22
kmeans, 4
length, big.matrix-method, 24
lm, 4
matrix, 2, 6
morder, 24
morderCols (morder), 24
mpermute (morder), 24
mpermuteCols (morder), 24
mwhich, 4, 26, 28
mwhich, big.matrix, ANY, ANY, ANY, character-method (mwhich-methods), 28
mwhich, big.matrix, ANY, ANY, ANY, missing-method (mwhich-methods), 28
mwhich, matrix, ANY, ANY, ANY, character-method (mwhich-methods), 28
mwhich, matrix, ANY, ANY, ANY, missing-method (mwhich-methods), 28
mwhich-methods, 28
ncol, big.matrix-method, 28
nrow, big.matrix-method
(ncol, big.matrix-method), 28
order, 25
print,big.matrix-method, 29
read.big.matrix, 4
read.big.matrix (write.big.matrix), 30
read.big.matrix, character-method (write.big.matrix), 30
shared. name (big.matrix), 6
shared.name,big.matrix-method (big.matrix), 6
split, 4
sub.big.matrix (is.sub.big.matrix), 22
sub.big.matrix,big.matrix-method
(is.sub.big.matrix), 22
sub.big.matrix,big.matrix.descriptor-method
(descriptor-class), 15
tail, big.matrix-method
(head, big.matrix-method), 21
typeof, big.matrix-method, 29
union, 26
which, 26-28
write.big.matrix, 30
write.big.matrix,big.matrix, character-method (write.big.matrix), 30

