# Package 'bigstatsr'

September 9, 2024

**Encoding** UTF-8

Type Package

Title Statistical Tools for Filebacked Big Matrices

Version 1.6.1

Date 2024-09-10

Description Easy-to-use, efficient, flexible and scalable statistical tools. Package bigstatsr provides and uses Filebacked Big Matrices via memory-mapping. It provides for instance matrix operations, Principal Component Analysis, sparse linear supervised models, utility functions and more <doi:10.1093/bioinformatics/bty185>.

License GPL-3

Language en-US

ByteCompile TRUE

**Depends** R (>= 3.4)

**Imports** bigassertr (>= 0.1.1), bigparallelr (>= 0.2.3), cowplot, foreach, ggplot2 (>= 3.0), graphics, methods, ps (>= 1.4), Rcpp, rmio (>= 0.4), RSpectra, stats, tibble, utils

LinkingTo Rcpp, RcppArmadillo, rmio

Suggests bigmemory (>= 4.5.33), bigreadr (>= 0.2), covr, data.table, dplyr, glmnet, hexbin, memuse, ModelMetrics, plotly, ppcor, RhpcBLASctl, spelling (>= 1.2), testthat

RoxygenNote 7.3.2

URL https://privefl.github.io/bigstatsr/

#### BugReports https://github.com/privefl/bigstatsr/issues

Collate 'AUC.R' 'FBM-attach.R' 'crochet.R' 'FBM.R' 'FBM-code256.R' 'FBM-copy.R' 'RcppExports.R' 'SVD.R' 'apply-parallelize.R' 'biglasso.R' 'bigstatsr-package.R' 'colstats.R' 'crossprodSelf.R' 'mult-mat.R' 'mult-vec.R' 'pcor.R' 'plot.R' 'predict.R' 'randomSVD.R' 'read-write.R' 'scaling.R' 'summary.R' 'tcrossprodSelf.R' 'transpose.R' 'univLinReg.R' 'univLogReg.R' 'utils-assert.R' 'utils.R' 'zzz.R'

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NeedsCompilation yes

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**Repository** CRAN

Date/Publication 2024-09-09 21:40:02 UTC

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asPlotlyText Plotly text

## Description

Convert a data.frame to plotly text

## Usage

asPlotlyText(df)

## Arguments

df A data.frame

## Value

A character vector of the length of df's number of rows.

## Examples

```
set.seed(1)
X <- big_attachExtdata()
svd <- big_SVD(X, big_scale(), k = 10)
p <- plot(svd, type = "scores")
pop <- rep(c("POP1", "POP2", "POP3"), c(143, 167, 207))
df <- data.frame(Population = pop, Index = 1:517)
plot(p2 <- p + ggplot2::aes(text = asPlotlyText(df)))
## Not run: plotly::ggplotly(p2, tooltip = "text")</pre>
```

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as\_scaling\_fun

#### Description

Convenience function to create a function to be used as parameter fun.scaling when you want to use your own precomputed center and scale.

## Usage

```
as_scaling_fun(center.col, scale.col, ind.col = seq_along(center.col))
```

#### Arguments

center.col	Vector of centers corresponding to ind.col.
<pre>scale.col</pre>	Vector of scales corresponding to ind.col.
ind.col	Column indices for which these are provided.

#### Value

A function to be used as parameter fun.scaling.

#### Examples

```
fun.scaling <- as_scaling_fun(1:6, 2:7)
fun.scaling(NULL, NULL, 1:3) # first two parameters X and ind.row are not used here
fun.scaling2 <- as_scaling_fun(1:6, 2:7, ind.col = 6:1)
fun.scaling2(NULL, NULL, 1:3)</pre>
```

```
X <- big_attachExtdata()
sc <- big_scale()(X)
fun <- as_scaling_fun(center = sc$center, scale = sc$scale)
obj.svd <- big_randomSVD(X, fun.scaling = fun)
obj.svd2 <- big_randomSVD(X, fun.scaling = big_scale())
all.equal(obj.svd, obj.svd2)</pre>
```

AUC

#### Description

AUC

Compute the Area Under the ROC Curve (AUC) of a predictor and possibly its 95% confidence interval.

## AUC

## Usage

```
AUC(pred, target, digits = NULL)
```

```
AUCBoot(pred, target, nboot = 10000, seed = NA, digits = NULL)
```

## Arguments

pred	Vector of predictions.
target	Vector of true labels (must have exactly two levels, no missing values).
digits	See round. Default doesn't use rounding.
nboot	Number of bootstrap samples used to evaluate the 95% CI. Default is 1e4.
seed	See set.seed. Use it for reproducibility. Default doesn't set any seed.

## Details

Other packages provide ways to compute the AUC (see this answer). I chose to compute the AUC through its statistical definition as a probability:

 $P(score(x_{case}) > score(x_{control})).$ 

Note that I consider equality between scores as a 50%-probability of one being greater than the other.

## Value

The AUC, a probability, and possibly its 2.5% and 97.5% quantiles (95% CI).

#### See Also

## wilcox.test

## Examples

```
set.seed(1)
```

```
AUC(c(0, 0), 0:1) # Equality of scores
AUC(c(0.2, 0.1, 1), c(0, 0, 1)) # Perfect AUC
x <- rnorm(100)
z <- rnorm(length(x), x, abs(x))
y <- as.numeric(z > 0)
print(AUC(x, y))
print(AUCBoot(x, y))
# Partial AUC
pAUC <- function(pred, target, p = 0.1) {
  val.min <- min(target)
  q <- quantile(pred[target == val.min], probs = 1 - p)
  ind <- (target != val.min) | (pred > q)
  bigstatsr::AUC(pred[ind], target[ind]) * p
}
```

```
pAUC(x, y)
pAUC(x, y, 0.2)
```

big\_apply

Split-Apply-Combine

## Description

A Split-Apply-Combine strategy to apply common R functions to a Filebacked Big Matrix.

## Usage

```
big_apply(
   X,
   a.FUN,
   a.combine = NULL,
   ind = cols_along(X),
   ncores = 1,
   block.size = block_size(nrow(X), ncores),
   ...
)
```

#### Arguments

Х	An object of class FBM.
a.FUN	The function to be applied to each subset matrix. It must take a Filebacked Big Matrix as first argument and ind, a vector of indices, which are used to split the data. For example, if you want to apply a function to X[ind.row, ind.col], you may use X[ind.row, ind.col[ind]] in a.FUN.
a.combine	Function to combine the results with do.call. This function should accept multiple arguments (). For example, you can use c, cbind, rbind. This package also provides function plus to add multiple arguments together. The default is NULL, in which case the results are not combined and are returned as a list, each element being the result of a block.
ind	Initial vector of subsetting indices. Default is the vector of all column indices.
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.
block.size	Maximum number of columns (or rows, depending on how you use ind for subsetting) read at once. Default uses block_size.
	Extra arguments to be passed to a. FUN.

## Details

This function splits indices in parts, then apply a given function to each subset matrix and finally combine the results. If parallelization is used, this function splits indices in parts for parallelization, then split again them on each core, apply a given function to each part and finally combine the results (on each cluster and then from each cluster). See also the corresponding vignette.

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#### big\_colstats

#### See Also

big\_parallelize bigparallelr::split\_parapply

#### Examples

```
X <- big_attachExtdata()</pre>
# get the means of each column
colMeans_sub <- function(X, ind) colMeans(X[, ind])</pre>
str(colmeans <- big_apply(X, a.FUN = colMeans_sub, a.combine = 'c'))</pre>
# get the norms of each column
colNorms_sub <- function(X, ind) sqrt(colSums(X[, ind]^2))</pre>
str(colnorms <- big_apply(X, colNorms_sub, a.combine = 'c'))</pre>
# get the sums of each row
# split along rows: need to change the "complete" `ind` parameter
str(rowsums <- big_apply(X, a.FUN = function(X, ind) rowSums(X[ind, ]),</pre>
                          ind = rows_along(X), a.combine = 'c',
                          block.size = 100))
# it is usually preferred to split along columns
# because matrices are stored by column.
str(rowsums2 <- big_apply(X, a.FUN = function(X, ind) rowSums(X[, ind]),</pre>
                           a.combine = 'plus'))
```

big\_colstats Standard univariate statistics

#### Description

Standard **univariate statistics** for columns of a Filebacked Big Matrix. For now, the sum and var are implemented (the mean and sd can easily be deduced, see examples).

#### Usage

big\_colstats(X, ind.row = rows\_along(X), ind.col = cols\_along(X), ncores = 1)

#### Arguments

Х	An object of class FBM.
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.

#### Value

Data.frame of two numeric vectors sum and var with the corresponding column statistics.

#### See Also

colSums apply

## Examples

set.seed(1)

```
X <- big_attachExtdata()</pre>
# Check the results
str(test <- big_colstats(X))</pre>
# Only with the first 100 rows
ind <- 1:100
str(test2 <- big_colstats(X, ind.row = ind))</pre>
plot(test$sum, test2$sum)
abline(lm(test2$sum ~ test$sum), col = "red", lwd = 2)
X.ind <- X[ind, ]
all.equal(test2$sum, colSums(X.ind))
all.equal(test2$var, apply(X.ind, 2, var))
# deduce mean and sd
# note that the are also implemented in big_scale()
means <- test2$sum / length(ind) # if using all rows,</pre>
                                   # divide by nrow(X) instead
all.equal(means, colMeans(X.ind))
sds <- sqrt(test2$var)</pre>
all.equal(sds, apply(X.ind, 2, sd))
```

big\_copy

Copy as a Filebacked Big Matrix

## Description

Deep copy of a Filebacked Big Matrix with possible subsetting. This should also work for any matrix-like object.

#### Usage

```
big_copy(
   X,
   ind.row = rows_along(X),
   ind.col = cols_along(X),
   type = typeof(X),
   backingfile = tempfile(tmpdir = getOption("FBM.dir")),
   block.size = block_size(length(ind.row)),
   is_read_only = FALSE
)
```

## big\_copy

## Arguments

Х	Could be any matrix-like object.
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
type	Type of the Filebacked Big Matrix (default is double). Either
	<ul> <li>"double" (double precision – 64 bits)</li> </ul>
	<ul> <li>"float" (single precision – 32 bits)</li> </ul>
	• "integer"
	• "unsigned short": can store integer values from 0 to 65535. It has vocation to become the basis for a FBM. code65536.
	• "raw" or "unsigned char": can store integer values from 0 to 255. It is the basis for class FBM.code256 in order to access 256 arbitrary different numeric values. It is used in package <b>bigsnpr</b> .
backingfile	Path to the file storing the FBM data on disk. An extension ".bk" will be automatically added. Default stores in the temporary directory, which you can change using global option "FBM.dir".
block.size	Maximum number of columns read at once. Default uses block_size.
is_read_only	Whether the FBM is read-only? Default is FALSE.

## Value

A copy of X as a new FBM object.

## Examples

```
X <- FBM(10, 10, init = 1:100)
X[]
X2 <- big_copy(X, ind.row = 1:5)
X2[]
mat <- matrix(101:200, 10)
X3 <- big_copy(mat, type = "double")  # as_FBM() would be faster here
X3[]
X.code <- big_attachExtdata()
class(X.code)
X2.code <- big_copy(X.code)
class(X2.code)
all.equal(X.code[], X2.code[])</pre>
```

big\_cor

## Description

Compute the (Pearson) correlation matrix of a Filebacked Big Matrix.

## Usage

```
big_cor(
   X,
   ind.row = rows_along(X),
   ind.col = cols_along(X),
   block.size = block_size(nrow(X)),
   backingfile = tempfile(tmpdir = getOption("FBM.dir"))
)
```

### Arguments

Х	An object of class FBM.
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
block.size	Maximum number of columns read at once. Default uses block_size.
backingfile	Path to the file storing the FBM data on disk. An extension ".bk" will be automatically added. Default stores in the temporary directory, which you can change using global option "FBM.dir".

## Value

A temporary FBM, with the following two attributes:

- a numeric vector center of column scaling,
- a numeric vector scale of column scaling.

#### Matrix parallelization

Large matrix computations are made block-wise and won't be parallelized in order to not have to reduce the size of these blocks. Instead, you can use the MKL or OpenBLAS in order to accelerate these block matrix computations. You can control the number of cores used by these optimized matrix libraries with bigparallelr::set\_blas\_ncores().

## See Also

cor big\_crossprodSelf

## big\_counts

## Examples

```
X <- FBM(13, 17, init = rnorm(221))
# Comparing with cor
K <- big_cor(X)
class(K)
dim(K)
K$backingfile
true <- cor(X[])
all.equal(K[], true)
# Using only half of the data
n <- nrow(X)
ind <- sort(sample(n, n/2))
K2 <- big_cor(X, ind.row = ind)
true2 <- cor(X[ind, ])
all.equal(K2[], true2)</pre>
```

big\_counts

Counts for class FBM.code256

## Description

Counts by columns (or rows) the number of each unique element of a FBM. code256.

## Usage

```
big_counts(
   X.code,
   ind.row = rows_along(X.code),
   ind.col = cols_along(X.code),
   byrow = FALSE
)
```

## Arguments

X.code	An object of class FBM.code256.
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
byrow	Count by rows rather than by columns? Default is FALSE (count by columns).

## Value

A matrix of counts of K x m (or n) elements, where

- K is the number of unique elements of the BM. code,
- n is its number of rows,
- m is its number of columns.

# Beware that K is up to 256. So, if you apply this on a Filebacked Big Matrix of one million columns, you will create a matrix of nearly 1GB!

## Examples

```
X <- big_attachExtdata()
class(X) # big_counts() is available for class FBM.code256 only
X[1:5, 1:8]
# by columns
big_counts(X, ind.row = 1:5, ind.col = 1:8)
# by rows
big_counts(X, ind.row = 1:5, ind.col = 1:8, byrow = TRUE)</pre>
```

big\_cprodMat Cross-product with a matrix

## Description

Cross-product between a Filebacked Big Matrix and a matrix.

#### Usage

```
big_cprodMat(
    X,
    A.row,
    ind.row = rows_along(X),
    ind.col = cols_along(X),
    ncores = 1,
    block.size = block_size(nrow(X), ncores),
    center = NULL,
    scale = NULL
)
## S4 method for signature 'FBM,matrix'
crossprod(x, y)
## S4 method for signature 'FBM,matrix'
```

## big\_cprodMat

```
tcrossprod(x, y)
## S4 method for signature 'matrix,FBM'
crossprod(x, y)
## S4 method for signature 'matrix,FBM'
tcrossprod(x, y)
```

## Arguments

<ul><li>A. row A matrix with length(ind.row) rows.</li><li>ind.row An optional vector of the row indices that are used. If not specified, all rows are used. Don't use negative indices.</li></ul>	Х	An object of class FBM.
	A.row	A matrix with length(ind.row) rows.
	ind.row	
ind.col An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>	ind.col	
ncores Number of cores used. Default doesn't use parallelism. You may use nb_cores.	ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.
block.size Maximum number of columns read at once. Default uses block_size.	block.size	Maximum number of columns read at once. Default uses block_size.
center Vector of same length of ind.col to subtract from columns of X.	center	Vector of same length of ind.col to subtract from columns of X.
scale Vector of same length of ind.col to divide from columns of X.	scale	Vector of same length of ind.col to divide from columns of X.
x A 'double' FBM or a matrix.	х	A 'double' FBM or a matrix.
y A 'double' FBM or a matrix.	У	A 'double' FBM or a matrix.

## Value

 $X^T \cdot A.$ 

## Matrix parallelization

Large matrix computations are made block-wise and won't be parallelized in order to not have to reduce the size of these blocks. Instead, you can use the MKL or OpenBLAS in order to accelerate these block matrix computations. You can control the number of cores used by these optimized matrix libraries with bigparallelr::set\_blas\_ncores().

## Examples

```
X <- big_attachExtdata()
n <- nrow(X)
m <- ncol(X)
A <- matrix(0, n, 10); A[] <- rnorm(length(A))
test <- big_cprodMat(X, A)
true <- crossprod(X[], A)
all.equal(test, true)
X2 <- big_copy(X, type = "double")
all.equal(crossprod(X2, A), true)</pre>
```

```
# subsetting
ind.row <- sample(n, n/2)
ind.col <- sample(m, m/2)
tryCatch(test2 <- big_cprodMat(X, A, ind.row, ind.col),
        error = function(e) print(e))
# returns an error. You need to use the subset of A:
test2 <- big_cprodMat(X, A[ind.row, ], ind.row, ind.col)
true2 <- crossprod(X[ind.row, ind.col], A[ind.row, ])
all.equal(test2, true2)
```

big\_cprodVec Cross-product with a vector

## Description

Cross-product between a Filebacked Big Matrix and a vector.

#### Usage

```
big_cprodVec(
   X,
   y.row,
   ind.row = rows_along(X),
   ind.col = cols_along(X),
   center = NULL,
   scale = NULL,
   ncores = 1
)
```

## Arguments

Х	An object of class FBM.
y.row	A vector of same size as ind.row.
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
center	Vector of same length of ind.col to subtract from columns of X.
scale	Vector of same length of ind.col to divide from columns of X.
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.

#### Value

 $X^T \cdot y.$ 

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## big\_crossprodSelf

## Examples

```
X <- big_attachExtdata()</pre>
n <- nrow(X)
m <- ncol(X)
y <- rnorm(n)</pre>
test <- big_cprodVec(X, y)</pre>
                                           # vector
true <- crossprod(X[], y) # one-column matrix</pre>
all.equal(test, as.numeric(true))
# subsetting
ind.row <- sample(n, n/2)</pre>
ind.col <- sample(m, m/2)</pre>
tryCatch(test2 <- big_cprodVec(X, y, ind.row, ind.col),</pre>
         error = function(e) print(e))
# returns an error. You need to use the subset of y:
test2 <- big_cprodVec(X, y[ind.row], ind.row, ind.col)</pre>
true2 <- crossprod(X[ind.row, ind.col], y[ind.row])</pre>
all.equal(test2, as.numeric(true2))
```

big\_crossprodSelf Crossprod

## Description

Compute  $X.row^T X.row$  for a Filebacked Big Matrix X after applying a particular scaling to it.

#### Usage

```
big_crossprodSelf(
    X,
    fun.scaling = big_scale(center = FALSE, scale = FALSE),
    ind.row = rows_along(X),
    ind.col = cols_along(X),
    block.size = block_size(nrow(X)),
    backingfile = tempfile(tmpdir = getOption("FBM.dir"))
)
## S4 method for signature 'FBM,missing'
crossprod(x, y)
```

#### Arguments

X An object of class FBM.

fun.scaling A function with parameters X, ind.row and ind.col, and that returns a data.frame with \$center and \$scale for the columns corresponding to ind.col, to scale each of their elements such as followed:

$$\frac{X_{i,j} - center_j}{scale_j}.$$

	Default doesn't use any scaling. You can also provide your own center and scale by using as_scaling_fun().
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
block.size	Maximum number of columns read at once. Default uses block_size.
backingfile	Path to the file storing the FBM data on disk. An extension ".bk" will be automatically added. Default stores in the temporary directory, which you can change using global option "FBM.dir".
x	A 'double' FBM.
У	Missing.

#### Value

A temporary FBM, with the following two attributes:

- a numeric vector center of column scaling,
- a numeric vector scale of column scaling.

## Matrix parallelization

Large matrix computations are made block-wise and won't be parallelized in order to not have to reduce the size of these blocks. Instead, you can use the MKL or OpenBLAS in order to accelerate these block matrix computations. You can control the number of cores used by these optimized matrix libraries with bigparallelr::set\_blas\_ncores().

#### See Also

crossprod

## Examples

```
X <- FBM(13, 17, init = rnorm(221))
true <- crossprod(X[])
# No scaling
K1 <- crossprod(X)
class(K1)</pre>
```

all.equal(K1, true)

K2 <- big\_crossprodSelf(X)</pre>

## big\_increment

```
class(K2)
K2$backingfile
all.equal(K2[], true)
# big_crossprodSelf() provides some scaling and subsetting
# Example using only half of the data:
n <- nrow(X)
ind <- sort(sample(n, n/2))
K3 <- big_crossprodSelf(X, fun.scaling = big_scale(), ind.row = ind)
true2 <- crossprod(scale(X[ind, ]))
all.equal(K3[], true2)</pre>
```

big\_increment Increment an FBM

## Description

Increment an FBM

#### Usage

big\_increment(X, add, use\_lock = FALSE)

## Arguments

Х	An FBM (of type double) to increment.
add	A matrix of same dimensions as X. Or a vector of same size.
use_lock	Whether to use locks when incrementing. Default is FALSE. This is useful when incrementing in parallel.

## Value

Returns nothing (NULL, invisibly).

## Examples

```
X <- FBM(10, 10, init = 0)
mat <- matrix(rnorm(100), 10, 10)
big_increment(X, mat)
all.equal(X[], mat)
big_increment(X, mat)
all.equal(X[], 2 * mat)</pre>
```

#### Description

A Split-Apply-Combine strategy to parallelize the evaluation of a function.

## Usage

```
big_parallelize(
   X,
   p.FUN,
   p.combine = NULL,
   ind = cols_along(X),
   ncores = nb_cores(),
   ...
)
```

## Arguments

Х	An object of class FBM.
p.FUN	The function to be applied to each subset matrix. It must take a Filebacked Big Matrix as first argument and ind, a vector of indices, which are used to split the data. For example, if you want to apply a function to X[ind.row, ind.col], you may use X[ind.row, ind.col[ind]] in a.FUN.
p.combine	Function to combine the results with do.call. This function should accept multiple arguments (). For example, you can use c, cbind, rbind. This package also provides function plus to add multiple arguments together. The default is NULL, in which case the results are not combined and are returned as a list, each element being the result of a block.
ind	Initial vector of subsetting indices. Default is the vector of all column indices.
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.
	Extra arguments to be passed to p. FUN.

#### Details

This function splits indices in parts, then apply a given function to each part and finally combine the results.

#### Value

Return a list of ncores elements, each element being the result of one of the cores, computed on a block. The elements of this list are then combined with do.call(p.combine, .) if p.combined is given.

#### big\_parallelize

#### See Also

big\_apply bigparallelr::split\_parapply

#### Examples

```
## Not run: # CRAN is super slow when parallelism.
 X <- big_attachExtdata()</pre>
 ### Computation on all the matrix
 true <- big_colstats(X)</pre>
 big_colstats_sub <- function(X, ind) {</pre>
   big_colstats(X, ind.col = ind)
 }
 # 1. the computation is split along all the columns
 # 2. for each part the computation is done, using `big_colstats`
 # 3. the results (data.frames) are combined via `rbind`.
 test <- big_parallelize(X, p.FUN = big_colstats_sub,</pre>
                           p.combine = 'rbind', ncores = 2)
 all.equal(test, true)
 ### Computation on a part of the matrix
 n <- nrow(X)
 m <- ncol(X)
 rows <- sort(sample(n, n/2)) # sort to provide some locality in accesses
 cols <- sort(sample(m, m/2)) # idem</pre>
 true2 <- big_colstats(X, ind.row = rows, ind.col = cols)</pre>
 big_colstats_sub2 <- function(X, ind, rows, cols) {</pre>
   big_colstats(X, ind.row = rows, ind.col = cols[ind])
 }
 # This doesn't work because, by default, the computation is spread
 # along all columns. We must explicitly specify the `ind` parameter.
 tryCatch(big_parallelize(X, p.FUN = big_colstats_sub2,
                           p.combine = 'rbind', ncores = 2,
                            rows = rows, cols = cols),
           error = function(e) message(e))
 # This now works, using `ind = seq_along(cols)`.
 test2 <- big_parallelize(X, p.FUN = big_colstats_sub2,</pre>
                            p.combine = 'rbind', ncores = 2,
                            ind = seq_along(cols),
                            rows = rows, cols = cols)
 all.equal(test2, true2)
```

## End(Not run)

big\_prodMat

## Description

Product between a Filebacked Big Matrix and a matrix.

## Usage

```
big_prodMat(
    X,
    A.col,
    ind.row = rows_along(X),
    ind.col = cols_along(X),
    ncores = 1,
    block.size = block_size(nrow(X), ncores),
    center = NULL,
    scale = NULL
)
## S4 method for signature 'FBM,matrix'
x %*% y
## S4 method for signature 'matrix,FBM'
x %*% y
```

## Arguments

Х	An object of class FBM.
A.col	A matrix with length(ind.col) rows.
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.
block.size	Maximum number of columns read at once. Default uses block_size.
center	Vector of same length of ind.col to subtract from columns of X.
scale	Vector of same length of ind.col to divide from columns of X.
x	A 'double' FBM or a matrix.
у	A 'double' FBM or a matrix.

#### Value

 $X\cdot A.$ 

## big\_prodVec

#### Matrix parallelization

Large matrix computations are made block-wise and won't be parallelized in order to not have to reduce the size of these blocks. Instead, you can use the MKL or OpenBLAS in order to accelerate these block matrix computations. You can control the number of cores used by these optimized matrix libraries with bigparallelr::set\_blas\_ncores().

## Examples

```
X <- big_attachExtdata()</pre>
n <- nrow(X)
m <- ncol(X)
A <- matrix(0, m, 10); A[] <- rnorm(length(A))</pre>
test <- big_prodMat(X, A)</pre>
true <- X[] %*% A
all.equal(test, true)
X2 <- big_copy(X, type = "double")</pre>
all.equal(X2 %*% A, true)
# subsetting
ind.row <- sample(n, n/2)</pre>
ind.col <- sample(m, m/2)</pre>
tryCatch(test2 <- big_prodMat(X, A, ind.row, ind.col),</pre>
          error = function(e) print(e))
# returns an error. You need to use the subset of A:
test2 <- big_prodMat(X, A[ind.col, ], ind.row, ind.col)</pre>
true2 <- X[ind.row, ind.col] %*% A[ind.col, ]</pre>
all.equal(test2, true2)
```

big\_prodVec

Product with a vector

#### Description

Product between a Filebacked Big Matrix and a vector.

#### Usage

```
big_prodVec(
    X,
    y.col,
    ind.row = rows_along(X),
    ind.col = cols_along(X),
    center = NULL,
    scale = NULL,
    ncores = 1
)
```

#### Arguments

Х	An object of class FBM.
y.col	A vector of same size as ind.col.
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
center	Vector of same length of ind.col to subtract from columns of X.
scale	Vector of same length of ind.col to divide from columns of X.
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.

## Value

 $X \cdot y$ .

## Examples

```
X <- big_attachExtdata()</pre>
n <- nrow(X)
m <- ncol(X)
y <- rnorm(m)
test <- big_prodVec(X, y)</pre>
                                  # vector
true <- X[] %*% y # one-column matrix</pre>
all.equal(test, as.numeric(true))
# subsetting
ind.row <- sample(n, n/2)</pre>
ind.col <- sample(m, m/2)</pre>
tryCatch(test2 <- big_prodVec(X, y, ind.row, ind.col),</pre>
         error = function(e) print(e))
# returns an error. You need to use the subset of y:
test2 <- big_prodVec(X, y[ind.col], ind.row, ind.col)</pre>
true2 <- X[ind.row, ind.col] %*% y[ind.col]</pre>
all.equal(test2, as.numeric(true2))
```

big\_randomSVD Randomized partial SVD

#### Description

An algorithm for partial SVD (or PCA) of a Filebacked Big Matrix based on the algorithm in RSpectra (by Yixuan Qiu and Jiali Mei).

This algorithm is linear in time in all dimensions and is very memory-efficient. Thus, it can be used on very large big.matrices.

## big\_randomSVD

## Usage

```
big_randomSVD(
   X,
   fun.scaling = big_scale(center = FALSE, scale = FALSE),
   ind.row = rows_along(X),
   ind.col = cols_along(X),
   k = 10,
   tol = 1e-04,
   verbose = FALSE,
   ncores = 1,
   fun.prod = big_prodVec,
   fun.cprod = big_cprodVec
)
```

## Arguments

Х	An object of class FBM.
fun.scaling	A function with parameters X, ind.row and ind.col, and that returns a data.frame with \$center and \$scale for the columns corresponding to ind.col, to scale each of their elements such as followed:
	$\frac{X_{i,j} - center_j}{scale_j}.$
	Default doesn't use any scaling. You can also provide your own center and scale by using as_scaling_fun().
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
k	Number of singular vectors/values to compute. Default is 10. This algorithm should be used to compute only a few singular vectors/values.
tol	Precision parameter of svds. Default is 1e-4.
verbose	Should some progress be printed? Default is FALSE.
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.
fun.prod	Function that takes 6 arguments (in this order):
	• a matrix-like object X,
	• a vector x,
	• a vector of row indices ind.row of X,
	• a vector of column indices ind.col of X,
	• a vector of column centers (corresponding to ind.col),
	• a vector of column scales (corresponding to ind.col), and compute the product of X (subsetted and scaled) with x.
fun.cprod	Same as fun.prod, but for the <i>transpose</i> of X.

## Value

A named list (an S3 class "big\_SVD") of

- d, the singular values,
- u, the left singular vectors,
- v, the right singular vectors,
- niter, the number of the iteration of the algorithm,
- nops, number of Matrix-Vector multiplications used,
- center, the centering vector,
- scale, the scaling vector.

Note that to obtain the Principal Components, you must use predict on the result. See examples.

#### Note

The idea of using this Implicitly Restarted Arnoldi Method algorithm comes from G. Abraham, Y. Qiu, and M. Inouye, FlashPCA2: principal component analysis of biobank-scale genotype datasets, bioRxiv: doi:10.1101/094714.

It proved to be faster than our implementation of the "blanczos" algorithm in Rokhlin, V., Szlam, A., & Tygert, M. (2010). A Randomized Algorithm for Principal Component Analysis. SIAM Journal on Matrix Analysis and Applications, 31(3), 1100-1124. doi:10.1137/080736417.

#### See Also

svds

#### Examples

```
set.seed(1)
X <- big_attachExtdata()</pre>
K <- 10
# Using only half of the data for "training"
n <- nrow(X)
ind <- sort(sample(n, n/2))</pre>
test <- big_randomSVD(X, fun.scaling = big_scale(), ind.row = ind, k = K)</pre>
str(test)
pca <- prcomp(X[ind, ], center = TRUE, scale. = TRUE)</pre>
# same scaling
all.equal(test$center, pca$center)
all.equal(test$scale, pca$scale)
# use this function to predict scores
class(test)
scores <- predict(test)</pre>
# scores and loadings are the same or opposite
```

## big\_read

```
plot(scores, pca$x[, 1:K])
plot(test$v, pca$rotation[, 1:K])
plot(test$u)
plot(test, type = "scores")
# projecting on new data
ind2 <- setdiff(rows_along(X), ind)
scores.test2 <- predict(test, X, ind.row = ind2)
scores.test3 <- predict(pca, X[-ind, ])
plot(scores.test2, scores.test3[, 1:K])</pre>
```

big\_read

Read a file as FBM

## Description

Read a file as a Filebacked Big Matrix by using package {bigreadr}. For a mini-tutorial, please see this vignette.

#### Usage

```
big_read(
  file,
  select,
  filter = NULL,
  type = c("double", "float", "integer", "unsigned short", "unsigned char", "raw"),
  backingfile = drop_ext(file),
   ...
)
```

#### Arguments

file	File to read.
select	Indices of columns to read (sorted). The length of select will be the number of columns of the resulting FBM.
filter	Vector used to subset the rows of each data frame.
type	Type of the Filebacked Big Matrix (default is double). Either
	<ul> <li>"double" (double precision – 64 bits)</li> </ul>
	<ul> <li>"float" (single precision – 32 bits)</li> </ul>
	• "integer"
	• "unsigned short": can store integer values from 0 to 65535. It has voca- tion to become the basis for a FBM.code65536.
	• "raw" or "unsigned char": can store integer values from 0 to 255. It is the basis for class FBM.code256 in order to access 256 arbitrary different numeric values. It is used in package <b>bigsnpr</b> .

Path to the file storing the FBM data on disk. An extension ".bk" will be auto- matically added. Default uses file without its extension.
Arguments passed on to bigreadr::big_fread2
nb_parts Number of parts in which to split reading (and transforming). Parts are referring to blocks of selected columns. Default uses part_size to set a good value.
skip Number of lines to skip at the beginning of file.
progress Show progress? Default is FALSE.
<code>part_size</code> Size of the parts if <code>nb_parts</code> is not supplied. Default is 500 $\star$ 1024^2 (500 MB).

## Value

A Filebacked Big Matrix of type type with length(select) columns.

big\_scale

Some scaling functions

## Description

Some scaling functions for a Filebacked Big Matrix to be used as the fun.scaling parameter of some functions of this package.

#### Usage

big\_scale(center = TRUE, scale = TRUE)

#### Arguments

center	A logical value: whether to return means or 0s.
scale	A logical value: whether to return standard deviations or 1s. You can't use scale without using center.

#### Details

One could think about less common scalings, such as for example the "y-aware" scaling which uses the inverse of betas of column-wise linear regression as scaling. See this post for details. It would be easy to implement it using big\_colstats to get column means and big\_univLinReg to get betas (and then inverse them).

#### Value

A new **function** that returns a data.frame of two vectors "center" and "scale" which are of the length of ind.col.

## big\_spLinReg

#### See Also

as\_scaling\_fun

## Examples

```
X <- big_attachExtdata()</pre>
```

```
# No scaling
big_noscale <- big_scale(center = FALSE, scale = FALSE)
class(big_noscale) # big_scale returns a new function
str(big_noscale(X))
big_noscale2 <- big_scale(center = FALSE)
str(big_noscale2(X)) # you can't scale without centering
# Centering
```

```
big_center <- big_scale(scale = FALSE)
str(big_center(X))
# + scaling
str(big_scale()(X))</pre>
```

big\_spLinReg Sparse linear regression

#### Description

Fit lasso (or elastic-net) penalized linear regression for a Filebacked Big Matrix. Covariables can be added (/!\ penalized by default /!\).

#### Usage

```
big_spLinReg(
 Χ,
 y.train,
  ind.train = rows_along(X),
  ind.col = cols_along(X),
  covar.train = NULL,
 base.train = NULL,
 pf.X = NULL,
  pf.covar = NULL,
  alphas = 1,
  power_scale = 1,
  power_adaptive = 0,
 K = 10,
  ind.sets = NULL,
  nlambda = 200,
  nlam.min = 50,
  n.abort = 10,
 dfmax = 50000,
```

```
warn = TRUE,
ncores = 1,
...
```

## Arguments

Х	An object of class FBM.
y.train	Vector of responses, corresponding to ind.train.
ind.train	An optional vector of the row indices that are used, for the training part. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
covar.train	Matrix of covariables to be added in each model to correct for confounders (e.g. the scores of PCA), corresponding to ind.train. Default is NULL and corresponds to only adding an intercept to each model. You can use covar_from_df() to convert from a data frame.
base.train	Vector of base predictions. Model will be learned starting from these predic- tions. This can be useful if you want to previously fit a model with large-effect variables that you don't want to penalize.
pf.X	A multiplicative factor for the penalty applied to each coefficient. If supplied, pf.X must be a numeric vector of the same length as ind.col. Default is all 1. The purpose of pf.X is to apply differential penalization if some coefficients are thought to be more likely than others to be in the model. Setting SOME to 0 allows to have unpenalized coefficients.
pf.covar	Same as pf.X, but for covar.train. You might want to set some to 0 as variables with large effects can mask small effects in penalized regression.
alphas	The elastic-net mixing parameter that controls the relative contribution from the lasso (11) and the ridge (12) penalty. The penalty is defined as
	$\alpha   \beta  _1 + (1 - \alpha)/2   \beta  _2^2.$
	alpha = 1 is the lasso penalty and alpha in between 0 (1e-4) and 1 is the elastic- net penalty. Default is 1. You can pass multiple values, and only one will be used (optimized by grid-search).
power_scale	When using lasso (alpha = 1), penalization to apply that is equivalent to scaling genotypes dividing by (standard deviation)^power_scale. Default is 1 and corresponding to standard scaling. Using 0 would correspond to using unscaled variables and using 0.5 is Pareto scaling. If you e.g. use power_scale = $c(0, 0.5, 1)$ , the best value in CMSA will be used (just like with alphas).
power_adaptive	Multiplicative penalty factor to apply to variables in the form of $1 / m_j^p$ ower_adaptive, where m_j is the marginal statistic for variable j. Default is 0, which effectively disables this option. If you e.g. use power_adaptive = c(0, 0.5, 1.5), the best value in CMSA will be used (just like with alphas).
К	Number of sets used in the Cross-Model Selection and Averaging (CMSA) pro- cedure. Default is 10.

ind.sets	Integer vectors of values between 1 and K specifying which set each index of the training set is in. Default randomly assigns these values but it can be useful to set this vector for reproducibility, or if you want to refine the grid-search over alphas using the same sets.
nlambda	The number of lambda values. Default is 200.
nlam.min	Minimum number of lambda values to investigate. Default is 50.
n.abort	Number of lambda values for which prediction on the validation set must de- crease before stopping. Default is 10.
dfmax	Upper bound for the number of nonzero coefficients. Default is 50e3 because, for large data sets, computational burden may be heavy for models with a large number of nonzero coefficients.
warn	Whether to warn if some models may not have reached a minimum. Default is TRUE.
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.
	Arguments passed on to COPY_biglasso_main
	lambda.min.ratio The smallest value for lambda, as a fraction of lambda.max. Default is .0001 if the number of observations is larger than the number of variables and .001 otherwise.
	eps Convergence threshold for inner coordinate descent. The algorithm iterates until the maximum change in the objective after any coefficient update is less than eps times the null deviance. Default value is 1e-5.
	max.iter Maximum number of iterations. Default is 1000. return.all Deprecated. Now always return all models.
	recurrent Depretated. Now always return an models.

#### Details

This is a modified version of one function of package biglasso. It adds the possibility to train models with covariables and use many types of FBM (not only double ones). Yet, it only corresponds to screen = "SSR" (Sequential Strong Rules).

Also, to remove the choice of the lambda parameter, we introduce the Cross-Model Selection and Averaging (CMSA) procedure:

- 1. This function separates the training set in K folds (e.g. 10).
- 2. In turn,
  - each fold is considered as an inner validation set and the others (K 1) folds form an inner training set,
  - the model is trained on the inner training set and the corresponding predictions (scores) for the inner validation set are computed,
  - the vector of scores which maximizes log-likelihood is determined,
  - the vector of coefficients corresponding to the previous vector of scores is chosen.
- 3. The K resulting vectors of coefficients are then averaged into one final vector of coefficients.

#### Value

Return an object of class  $big_sp_list$  (a list of length(alphas) x K) that has 3 methods predict, summary and plot.

#### References

Tibshirani, R., Bien, J., Friedman, J., Hastie, T., Simon, N., Taylor, J. and Tibshirani, R. J. (2012), Strong rules for discarding predictors in lasso-type problems. Journal of the Royal Statistical Society: Series B (Statistical Methodology), 74: 245-266. doi:10.1111/j.14679868.2011.01004.x.

Zeng, Y., and Breheny, P. (2017). The biglasso Package: A Memory- and Computation-Efficient Solver for Lasso Model Fitting with Big Data in R. doi:10.32614/RJ2021001.

Privé, F., Aschard, H., and Blum, M. G.B. (2019). Efficient implementation of penalized regression for genetic risk prediction. Genetics, 212: 65-74. doi:10.1534/genetics.119.302019.

#### See Also

glmnet

#### Examples

```
set.seed(1)
# simulating some data
N <- 230
M <- 730
X <- FBM(N, M, init = rnorm(N * M, sd = 5))
y <- rowSums(X[, 1:10]) + rnorm(N)</pre>
covar <- matrix(rnorm(N * 3), N)</pre>
ind.train <- sort(sample(nrow(X), 150))</pre>
ind.test <- setdiff(rows_along(X), ind.train)</pre>
# fitting model for multiple lambdas and alphas
test <- big_spLinReg(X, y[ind.train], ind.train = ind.train,</pre>
                      covar.train = covar[ind.train, ],
                      alphas = c(1, 0.1), K = 3, warn = FALSE)
# peek at the models
plot(test)
summary(test, sort = TRUE)
summary(test, sort = TRUE)$message
# prediction for other data -> only the best alpha is used
summary(test, best.only = TRUE)
pred <- predict(test, X, ind.row = ind.test, covar.row = covar[ind.test, ])</pre>
plot(pred, y[ind.test], pch = 20); abline(0, 1, col = "red")
```

big\_spLogReg Sparse logistic regression

#### Description

Fit lasso (or elastic-net) penalized logistic regression for a Filebacked Big Matrix. Covariables can be added (/!\ penalized by default /!\).

## big\_spLogReg

## Usage

```
big_spLogReg(
 Χ,
 y01.train,
  ind.train = rows_along(X),
  ind.col = cols_along(X),
  covar.train = NULL,
 base.train = NULL,
 pf.X = NULL,
 pf.covar = NULL,
 alphas = 1,
 power_scale = 1,
 power_adaptive = 0,
 K = 10,
  ind.sets = NULL,
 nlambda = 200,
 nlam.min = 50,
 n.abort = 10,
 dfmax = 50000,
 warn = TRUE,
 ncores = 1,
  . . .
)
```

## Arguments

Х	An object of class FBM.
y01.train	Vector of responses, corresponding to ind.train. Must be only 0s and 1s.
ind.train	An optional vector of the row indices that are used, for the training part. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
covar.train	Matrix of covariables to be added in each model to correct for confounders (e.g. the scores of PCA), corresponding to ind.train. Default is NULL and corresponds to only adding an intercept to each model. You can use covar_from_df() to convert from a data frame.
base.train	Vector of base predictions. Model will be learned starting from these predic- tions. This can be useful if you want to previously fit a model with large-effect variables that you don't want to penalize.
pf.X	A multiplicative factor for the penalty applied to each coefficient. If supplied, $pf.X$ must be a numeric vector of the same length as ind.col. Default is all 1. The purpose of $pf.X$ is to apply differential penalization if some coefficients are thought to be more likely than others to be in the model. Setting SOME to 0 allows to have unpenalized coefficients.
pf.covar	Same as pf.X, but for covar.train. You might want to set some to 0 as variables with large effects can mask small effects in penalized regression.

alphas	The elastic-net mixing parameter that controls the relative contribution from the lasso (11) and the ridge (12) penalty. The penalty is defined as
	$\alpha   \beta  _1 + (1 - \alpha)/2   \beta  _2^2.$
	alpha = 1 is the lasso penalty and alpha in between $0$ (1e-4) and 1 is the elastic- net penalty. Default is 1. You can pass multiple values, and only one will be used (optimized by grid-search).
power_scale	When using lasso (alpha = 1), penalization to apply that is equivalent to scaling genotypes dividing by (standard deviation)^power_scale. Default is 1 and corresponding to standard scaling. Using 0 would correspond to using unscaled variables and using 0.5 is Pareto scaling. If you e.g. use power_scale = $c(0, 0.5, 1)$ , the best value in CMSA will be used (just like with alphas).
power_adaptive	Multiplicative penalty factor to apply to variables in the form of $1/m_j^p$ ower_adaptive, where m_j is the marginal statistic for variable j. Default is 0, which effectively disables this option. If you e.g. use power_adaptive = c(0, 0.5, 1.5), the best value in CMSA will be used (just like with alphas).
К	Number of sets used in the Cross-Model Selection and Averaging (CMSA) pro- cedure. Default is 10.
ind.sets	Integer vectors of values between 1 and K specifying which set each index of the training set is in. Default randomly assigns these values but it can be useful to set this vector for reproducibility, or if you want to refine the grid-search over alphas using the same sets.
nlambda	The number of lambda values. Default is 200.
nlam.min	Minimum number of lambda values to investigate. Default is 50.
n.abort	Number of lambda values for which prediction on the validation set must decrease before stopping. Default is 10.
dfmax	Upper bound for the number of nonzero coefficients. Default is 50e3 because, for large data sets, computational burden may be heavy for models with a large number of nonzero coefficients.
warn	Whether to warn if some models may not have reached a minimum. Default is TRUE.
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.
	Arguments passed on to COPY_biglasso_main
	<pre>lambda.min.ratio The smallest value for lambda, as a fraction of lambda.max. Default is .0001 if the number of observations is larger than the number of variables and .001 otherwise.</pre>
	eps Convergence threshold for inner coordinate descent. The algorithm iterates until the maximum change in the objective after any coefficient update is less than eps times the null deviance. Default value is 1e-5.
	max.iter Maximum number of iterations. Default is 1000.
	return.all Deprecated. Now always return all models.

#### Details

This is a modified version of one function of package biglasso. It adds the possibility to train models with covariables and use many types of FBM (not only double ones). Yet, it only corresponds to screen = "SSR" (Sequential Strong Rules).

Also, to remove the choice of the lambda parameter, we introduce the Cross-Model Selection and Averaging (CMSA) procedure:

- 1. This function separates the training set in K folds (e.g. 10).
- 2. In turn,
  - each fold is considered as an inner validation set and the others (K 1) folds form an inner training set,
  - the model is trained on the inner training set and the corresponding predictions (scores) for the inner validation set are computed,
  - the vector of scores which maximizes log-likelihood is determined,
  - the vector of coefficients corresponding to the previous vector of scores is chosen.
- 3. The K resulting vectors of coefficients are then averaged into one final vector of coefficients.

## Value

Return an object of class big\_sp\_list (a list of length(alphas) x K) that has 3 methods predict, summary and plot.

## References

Tibshirani, R., Bien, J., Friedman, J., Hastie, T., Simon, N., Taylor, J. and Tibshirani, R. J. (2012), Strong rules for discarding predictors in lasso-type problems. Journal of the Royal Statistical Society: Series B (Statistical Methodology), 74: 245-266. doi:10.1111/j.14679868.2011.01004.x.

Zeng, Y., and Breheny, P. (2017). The biglasso Package: A Memory- and Computation-Efficient Solver for Lasso Model Fitting with Big Data in R. doi:10.32614/RJ2021001.

Privé, F., Aschard, H., and Blum, M. G.B. (2019). Efficient implementation of penalized regression for genetic risk prediction. Genetics, 212: 65-74. doi:10.1534/genetics.119.302019.

#### See Also

glmnet

set.seed(2)

#### Examples

```
# simulating some data
N <- 230
M <- 730
X <- FBM(N, M, init = rnorm(N * M, sd = 5))
y01 <- as.numeric((rowSums(X[, 1:10]) + 2 * rnorm(N)) > 0)
covar <- matrix(rnorm(N * 3), N)
ind.train <- sort(sample(nrow(X), 150))</pre>
```

```
ind.test <- setdiff(rows_along(X), ind.train)</pre>
# fitting model for multiple lambdas and alphas
test <- big_spLogReg(X, y01[ind.train], ind.train = ind.train,</pre>
                     covar.train = covar[ind.train, ],
                     alphas = c(1, 0.1), K = 3, warn = FALSE)
# peek at the models
plot(test)
summary(test, sort = TRUE)
summary(test, sort = TRUE)$message
# prediction for other data -> only the best alpha is used
summary(test, best.only = TRUE)
pred <- predict(test, X, ind.row = ind.test, covar.row = covar[ind.test, ])</pre>
AUC(pred, y01[ind.test])
library(ggplot2)
qplot(pred, fill = as.logical(y01[ind.test]),
      geom = "density", alpha = I(0.4)) +
 labs(fill = "Case?") +
 theme_bigstatsr() +
 theme(legend.position = c(0.52, 0.8))
```

big\_SVD

Partial SVD

#### Description

An algorithm for partial SVD (or PCA) of a Filebacked Big Matrix through the eigen decomposition of the covariance between variables (primal) or observations (dual). Use this algorithm only if there is one dimension that is much smaller than the other. Otherwise use big\_randomSVD.

#### Usage

```
big_SVD(
   X,
   fun.scaling = big_scale(center = FALSE, scale = FALSE),
   ind.row = rows_along(X),
   ind.col = cols_along(X),
   k = 10,
   block.size = block_size(nrow(X))
)
```

```
)
```

#### Arguments

X An object of class FBM.

fun.scaling A function with parameters X, ind.row and ind.col, and that returns a data.frame with \$center and \$scale for the columns corresponding to ind.col, to scale each of their elements such as followed:

$$\frac{X_{i,j} - center_j}{scale_j}.$$

	Default doesn't use any scaling. You can also provide your own center and scale by using as_scaling_fun().
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
k	Number of singular vectors/values to compute. Default is 10. <b>This algorithm should be used to compute only a few singular vectors/values.</b> If more is needed, have a look at https://stackoverflow.com/a/46380540/6103040.
block.size	Maximum number of columns read at once. Default uses block_size.

#### Details

To get  $X = U \cdot D \cdot V^T$ ,

- if the number of observations is small, this function computes  $K_{(2)} = X \cdot X^T \approx U \cdot D^2 \cdot U^T$ and then  $V = X^T \cdot U \cdot D^{-1}$ ,
- if the number of variable is small, this function computes  $K_{(1)} = X^T \cdot X \approx V \cdot D^2 \cdot V^T$  and then  $U = X \cdot V \cdot D^{-1}$ ,
- if both dimensions are large, use big\_randomSVD instead.

#### Value

A named list (an S3 class "big\_SVD") of

- d, the singular values,
- u, the left singular vectors,
- v, the right singular vectors,
- center, the centering vector,
- scale, the scaling vector.

Note that to obtain the Principal Components, you must use predict on the result. See examples.

#### Matrix parallelization

Large matrix computations are made block-wise and won't be parallelized in order to not have to reduce the size of these blocks. Instead, you can use the MKL or OpenBLAS in order to accelerate these block matrix computations. You can control the number of cores used by these optimized matrix libraries with bigparallelr::set\_blas\_ncores().

#### See Also

prcomp

#### Examples

```
set.seed(1)
X <- big_attachExtdata()</pre>
n <- nrow(X)
# Using only half of the data
ind <- sort(sample(n, n/2))</pre>
test <- big_SVD(X, fun.scaling = big_scale(), ind.row = ind)</pre>
str(test)
plot(test$u)
pca <- prcomp(X[ind, ], center = TRUE, scale. = TRUE)</pre>
# same scaling
all.equal(test$center, pca$center)
all.equal(test$scale, pca$scale)
# scores and loadings are the same or opposite
# except for last eigenvalue which is equal to 0
# due to centering of columns
scores <- test$u %*% diag(test$d)</pre>
class(test)
scores2 <- predict(test) # use this function to predict scores</pre>
all.equal(scores, scores2)
dim(scores)
dim(pca$x)
tail(pca$sdev)
plot(scores2, pca$x[, 1:ncol(scores2)])
plot(test$v[1:100, ], pca$rotation[1:100, 1:ncol(scores2)])
# projecting on new data
X2 <- sweep(sweep(X[-ind, ], 2, test$center, '-'), 2, test$scale, '/')</pre>
scores.test <- X2 %*% test$v</pre>
ind2 <- setdiff(rows_along(X), ind)</pre>
scores.test2 <- predict(test, X, ind.row = ind2) # use this</pre>
all.equal(scores.test, scores.test2)
scores.test3 <- predict(pca, X[-ind, ])</pre>
plot(scores.test2, scores.test3[, 1:ncol(scores.test2)])
```

big\_tcrossprodSelf Tcrossprod

#### Description

Compute  $X.row X.row^T$  for a Filebacked Big Matrix X after applying a particular scaling to it.

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## big\_tcrossprodSelf

## Usage

```
big_tcrossprodSelf(
   X,
   fun.scaling = big_scale(center = FALSE, scale = FALSE),
   ind.row = rows_along(X),
   ind.col = cols_along(X),
   block.size = block_size(nrow(X))
)
## S4 method for signature 'FBM,missing'
```

## tcrossprod(x, y)

## Arguments

Х	An object of class FBM.
fun.scaling	A function with parameters X, ind.row and ind.col, and that returns a data.frame with \$center and \$scale for the columns corresponding to ind.col, to scale each of their elements such as followed:
	$\frac{X_{i,j} - center_j}{scale_j}.$
	Default doesn't use any scaling. You can also provide your own center and scale by using as_scaling_fun().
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
block.size	Maximum number of columns read at once. Default uses block_size.
х	A 'double' FBM.
У	Missing.

## Value

A temporary FBM, with the following two attributes:

- a numeric vector center of column scaling,
- a numeric vector scale of column scaling.

#### Matrix parallelization

Large matrix computations are made block-wise and won't be parallelized in order to not have to reduce the size of these blocks. Instead, you can use the MKL or OpenBLAS in order to accelerate these block matrix computations. You can control the number of cores used by these optimized matrix libraries with bigparallelr::set\_blas\_ncores().

# See Also

tcrossprod

#### Examples

```
X <- FBM(13, 17, init = rnorm(221))
true <- tcrossprod(X[])</pre>
# No scaling
K1 <- tcrossprod(X)</pre>
class(K1)
all.equal(K1, true)
K2 <- big_tcrossprodSelf(X)</pre>
class(K2)
K2$backingfile
all.equal(K2[], true)
# big_tcrossprodSelf() provides some scaling and subsetting
# Example using only half of the data:
n <- nrow(X)
ind <- sort(sample(n, n/2))</pre>
K3 <- big_tcrossprodSelf(X, fun.scaling = big_scale(), ind.row = ind)</pre>
true2 <- tcrossprod(scale(X[ind, ]))</pre>
all.equal(K3[], true2)
```

big\_transpose Transpose an FBM

## Description

This function implements a simple cache-oblivious algorithm for the transposition of a Filebacked Big Matrix.

#### Usage

```
big_transpose(X, backingfile = tempfile(tmpdir = getOption("FBM.dir")))
```

#### Arguments

Х	An object of class FBM.
backingfile	Path to the file storing the FBM data on disk. An extension ".bk" will be
	automatically added. Default stores in the temporary directory, which you can
	change using global option "FBM.dir".

#### Value

The new transposed FBM. Dimensions and type are automatically determined from the input FBM.

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## big\_univLinReg

## Examples

```
X <- FBM(6, 5, init = rnorm(30))
X[]
Xt <- big_transpose(X)
identical(t(X[]), Xt[])</pre>
```

big\_univLinReg Column-wise linear regression

## Description

Slopes of column-wise linear regressions of each column of a Filebacked Big Matrix, with some other associated statistics. Covariates can be added to correct for confounders.

## Usage

```
big_univLinReg(
   X,
   y.train,
   ind.train = rows_along(X),
   ind.col = cols_along(X),
   covar.train = NULL,
   thr.eigval = 1e-04,
   ncores = 1
)
```

# Arguments

Х	An object of class FBM.
y.train	Vector of responses, corresponding to ind.train.
ind.train	An optional vector of the row indices that are used, for the training part. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
covar.train	Matrix of covariables to be added in each model to correct for confounders (e.g. the scores of PCA), corresponding to ind.train. Default is NULL and corresponds to only adding an intercept to each model. You can use covar_from_df() to convert from a data frame.
thr.eigval	Threshold to remove "insignificant" singular vectors. Default is 1e-4.
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.

A data.frame with 3 elements:

- 1. the slopes of each regression,
- 2. the standard errors of each slope,
- 3. the t-scores associated with each slope. This is also an object of class mhtest. See methods(class = "mhtest").

#### See Also

lm

```
set.seed(1)
X <- big_attachExtdata()</pre>
n <- nrow(X)
y <- rnorm(n)
covar <- matrix(rnorm(n * 3), n)</pre>
X1 <- X[, 1] # only first column of the Filebacked Big Matrix
# Without covar
test <- big_univLinReg(X, y)</pre>
## New class `mhtest`
class(test)
attr(test, "transfo")
attr(test, "predict")
## plot results
plot(test)
plot(test, type = "Volcano")
## To get p-values associated with the test
test$p.value <- predict(test, log10 = FALSE)</pre>
str(test)
summary(lm(y ~ X1))$coefficients[2, ]
# With all data
str(big_univLinReg(X, y, covar = covar))
summary(lm(y ~ X1 + covar))$coefficients[2, ]
# With only half of the data
ind.train <- sort(sample(n, n/2))</pre>
str(big_univLinReg(X, y[ind.train],
                   covar.train = covar[ind.train, ],
                   ind.train = ind.train))
summary(lm(y ~ X1 + covar, subset = ind.train))$coefficients[2, ]
```

#### Description

Slopes of column-wise logistic regressions of each column of a Filebacked Big Matrix, with some other associated statistics. Covariates can be added to correct for confounders.

## Usage

```
big_univLogReg(
   X,
   y01.train,
   ind.train = rows_along(X),
   ind.col = cols_along(X),
   covar.train = NULL,
   tol = 1e-08,
   maxiter = 20,
   ncores = 1
)
```

#### Arguments

Х	An object of class FBM.
y01.train	Vector of responses, corresponding to ind.train. Must be only 0s and 1s.
ind.train	An optional vector of the row indices that are used, for the training part. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
covar.train	Matrix of covariables to be added in each model to correct for confounders (e.g. the scores of PCA), corresponding to ind.train. Default is NULL and corresponds to only adding an intercept to each model. You can use covar_from_df() to convert from a data frame.
tol	Relative tolerance to assess convergence of the coefficient. Default is 1e-8.
maxiter	Maximum number of iterations before giving up. Default is 20. Usually, convergence is reached within 3 or 4 iterations. If there is not convergence, glm is used instead for the corresponding column.
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.

## Details

If convergence is not reached by the main algorithm for some columns, the corresponding niter element is set to NA and a message is given. Then, glm is used instead for the corresponding column. If it can't converge either, all corresponding estimations are set to NA.

#### Value

A data.frame with 4 elements:

- 1. the slopes of each regression,
- 2. the standard errors of each slope,
- 3. the number of iteration for each slope. If is NA, this means that the algorithm didn't converge, and glm was used instead.
- 4. the z-scores associated with each slope. This is also an object of class mhtest. See methods(class = "mhtest").

## See Also

glm

#### Examples

set.seed(1)

```
X <- big_attachExtdata()
n <- nrow(X)
y01 <- sample(0:1, size = n, replace = TRUE)
covar <- matrix(rnorm(n * 3), n)</pre>
```

X1 <- X[, 1] # only first column of the Filebacked Big Matrix

```
# Without covar
test <- big_univLogReg(X, y01)</pre>
## new class `mhtest`
class(test)
attr(test, "transfo")
attr(test, "predict")
## plot results
plot(test)
plot(test, type = "Volcano")
## To get p-values associated with the test
test$p.value <- predict(test, log10 = FALSE)</pre>
str(test)
summary(glm(y01 ~ X1, family = "binomial"))$coefficients[2, ]
# With all data
str(big_univLogReg(X, y01, covar.train = covar))
summary(glm(y01 ~ X1 + covar, family = "binomial"))$coefficients[2, ]
# With only half of the data
ind.train <- sort(sample(n, n/2))</pre>
str(big_univLogReg(X, y01[ind.train],
                   covar.train = covar[ind.train, ],
                   ind.train = ind.train))
summary(glm(y01 ~ X1 + covar, family = "binomial",
            subset = ind.train))$coefficients[2, ]
```

big\_write

# Description

Write a file from a Filebacked Big Matrix (by parts).

## Usage

```
big_write(
   X,
   file,
   every_nrow,
   ...,
   ind.row = rows_along(X),
   ind.col = cols_along(X),
   progress = FALSE
)
```

## Arguments

Х	An object of class FBM.
file	File to write to.
every_nrow	Number of rows to write at once.
	Other arguments to be passed to data.table::fwrite, except x, file, append, row.names, col.names and showProgress.
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
progress	Show progress? Default is FALSE.

## Value

Input parameter file, invisibly.

```
X <- big_attachExtdata()
csv <- big_write(X, tempfile(), every_nrow = 100, progress = interactive())</pre>
```

block\_size

#### Description

It determines the value of block.size such that a matrix of doubles of size n x block.size takes less memory than getOption("bigstatsr.block.sizeGB") GigaBytes (default is 1GB).

## Usage

block\_size(n, ncores = 1)

#### Arguments

n	The number of rows.
ncores	The number of cores.

#### Value

An integer >= 1.

## Examples

```
block_size(1e3)
block_size(1e6)
block_size(1e6, 6)
```

covar\_from\_df Numeric matrix from data frame

## Description

Transform a data frame to a numeric matrix by one-hot encoding factors. The last factor value is always omitted to prevent having a singular matrix when adding a column of 1s (intercept) in models.

## Usage

covar\_from\_df(df)

## Arguments

df A data frame.

## Value

A numeric matrix.

## FBM-class

#### Examples

```
mat <- covar_from_df(iris)
head(mat)</pre>
```

FBM-class

Class FBM

## Description

A reference class for storing and accessing matrix-like data stored in files on disk. This is very similar to Filebacked Big Matrices provided by the **bigmemory** package (see the corresponding vignette).

Convert a matrix (or a data frame) to an FBM.

## Usage

```
FBM(
    nrow,
    ncol,
    type = c("double", "float", "integer", "unsigned short", "unsigned char", "raw"),
    init = NULL,
    backingfile = tempfile(tmpdir = getOption("FBM.dir")),
    create_bk = TRUE,
    is_read_only = FALSE
)
as_FBM(
    x,
    type = c("double", "float", "integer", "unsigned short", "unsigned char", "raw"),
    backingfile = tempfile(tmpdir = getOption("FBM.dir")),
    is_read_only = FALSE
)
```

#### Arguments

nrow	Number of rows.
ncol	Number of columns.
type	Type of the Filebacked Big Matrix (default is double). Either
	<ul> <li>"double" (double precision – 64 bits)</li> </ul>
	<ul> <li>"float" (single precision – 32 bits)</li> </ul>
	• "integer"
	• "unsigned short": can store integer values from 0 to 65535. It has voca- tion to become the basis for a FBM. code65536.
	• "raw" or "unsigned char": can store integer values from 0 to 255. It is the basis for class FBM.code256 in order to access 256 arbitrary different numeric values. It is used in package <b>bigsnpr</b> .

init	Either a single value (e.g. 0) or as many value as the number of elements of the FBM. <b>Default doesn't initialize the matrix.</b>
backingfile	Path to the file storing the FBM data on disk. An extension ".bk" will be automatically added. Default stores in the temporary directory, which you can change using global option "FBM.dir".
create_bk	Whether to create a backingfile (the default) or use an existing one (which should be named by the backingfile parameter and have an extension ".bk"). For example, this could be used to convert a filebacked big.matrix from package <b>bigmemory</b> to a FBM (see the corresponding vignette).
is_read_only	Whether the FBM is read-only? Default is FALSE.
х	A matrix or an data frame (2-dimensional data).

#### Details

An object of class FBM has many fields:

- \$address: address of the external pointer containing the underlying C++ object for read-only mapping, to be used as a XPtr<FBM> in C++ code
- \$extptr: (internal) use \$address instead
- \$address\_rw: address of the external pointer containing the underlying C++ object for read/write mapping, to be used as a XPtr<FBM\_RW> in C++ code
- \$extptr\_rw: (internal) use \$address\_rw instead
- \$nrow: number of rows
- \$ncol: number of columns
- \$type: (internal) use type\_size or type\_chr instead
- \$type\_chr: FBM type as character, e.g. "double"
- \$type\_size: size of FBM type in bytes (e.g. "double" is 8 and "float" is 4)
- \$backingfile or \$bk: File with extension 'bk' that stores the numeric data of the FBM
- \$rds: 'rds' file (that may not exist) corresponding to the 'bk' file
- \$is\_saved: whether this object is stored in \$rds?
- \$is\_read\_only: whether it is (not) allowed to modify data?

And some methods:

- \$save(): Save the FBM object in \$rds. Returns the FBM.
- add\_columns(<ncol\_add>): Add some columns to the FBM by appending the backingfile with some data. Returns the FBM invisibly.
- \$bm(): Get this object as a filebacked.big.matrix to be used by package {bigmemory}.
- \$bm.desc(): Get this object as a filebacked.big.matrix descriptor to be used by package {bigmemory}.
- \$check\_write\_permissions(): Error if the FBM is read-only.

#### See Also

big\_attach big\_copy

## FBM-methods

#### Examples

```
mat <- matrix(1:4, 2)</pre>
X_from_mat <- as_FBM(mat)</pre>
## You can save this object in an .rds file to use it in another session
X_from_mat$is_saved
X_from_mat$save()
X_from_mat$is_saved
(rds <- X_from_mat$rds)</pre>
## Use big_attach() to load the FBM object in another session
X_from_mat <- big_attach(rds)</pre>
## Standard accessors
X <- FBM(10, 10)
typeof(X)
X[] <- rnorm(length(X))</pre>
X[, 1:6]
X[] <- 1:100
X[, 1]
X[, -1]
X[, c(TRUE, FALSE)]
X[cbind(1:10, 1:10)] <- NA_real_
X[] # access as standard R matrix
X <- FBM(150, 5)
X[] <- iris ## you can replace with a df (but factors -> integers)
X2 <- as_FBM(iris)
identical(X[], X2[])
```

```
FBM-methods
```

Methods for the FBM class

#### Description

Methods for the FBM class

Accessor methods for class FBM. You can use positive and negative indices, logical indices (that are recycled) and also a matrix of indices (but only positive ones).

Dimension and type methods for class FBM.

#### Usage

```
## S4 method for signature 'FBM,ANY,ANY'
x[i, j, ..., drop = TRUE]
## S4 replacement method for signature 'FBM,ANY,ANY,ANY'
```

```
x[i, j, ...] <- value
## S4 method for signature 'FBM'
dim(x)
## S4 method for signature 'FBM'
length(x)
## S4 method for signature 'FBM'
typeof(x)
## S4 method for signature 'FBM'
diag(x)</pre>
```

## Arguments

х	A FBM object.
i	A vector of indices (or nothing). You can use positive and negative indices, logical indices (that are recycled) and also a matrix of indices (but only positive ones).
j	A vector of indices (or nothing). You can use positive and negative indices, logical indices (that are recycled).
	Not used. Just to make nargs work.
drop	Whether to delete the dimensions of a matrix which have one dimension equals to 1.
value	The values to replace. Should be of length 1 or of the same length of the subset to replace.
Value	

FBM.code256-class Class FBM.code256

## Description

A reference class for storing and accessing up to 256 arbitrary different values using a Filebacked Big Matrix of type unsigned char. Compared to a Filebacked Big Matrix, it adds a slot code which is used as a lookup table of size 256.

## Usage

```
FBM.code256(
    nrow,
    ncol,
    code = rep(NA_real_, 256),
    init = NULL,
    backingfile = tempfile(tmpdir = getOption("FBM.dir")),
    create_bk = TRUE,
```

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is\_read\_only = FALSE
)

add\_code256(x, code)

# Arguments

nrow	Number of rows.
ncol	Number of columns.
code	A numeric vector (of length 256). You should construct it with rep(NA_real_, 256) and then replace the values which are of interest to you.
init	Either a single value (e.g. 0) or as many value as the number of elements of the FBM. <b>Default doesn't initialize the matrix.</b>
backingfile	Path to the file storing the FBM data on disk. An extension ".bk" will be automatically added. Default stores in the temporary directory, which you can change using global option "FBM.dir".
create_bk	Whether to create a backingfile (the default) or use an existing one (which should be named by the backingfile parameter and have an extension ".bk"). For example, this could be used to convert a filebacked big.matrix from package <b>bigmemory</b> to a FBM (see the corresponding vignette).
is_read_only	Whether the FBM is read-only? Default is FALSE.
х	A FBM.

```
X <- FBM(10, 10, type = "raw")
X[] <- sample(as.raw(0:3), size = length(X), replace = TRUE)
X[]
# From an FBM of type 'raw' ('unsigned char')
code <- rep(NA_real_, 256)
code[1:3] <- c(1, 3, 5)
X.code <- add_code256(X, code)
X.code[]
# Or directly
X.code2 <- FBM.code256(10, 10, code, init = sample(as.raw(0:3), 100, TRUE))
X.code2[]
# Get a new FBM.code256 object with another code (but same underlying data)
X.code3 <- X.code$copy(code = rnorm(256))
all.equal(X.code$code256, code)</pre>
```

get\_beta

## Description

Combine sets of coefficients

## Usage

```
get_beta(betas, method = c("geometric-median", "mean-wise", "median-wise"))
```

## Arguments

betas	Matrix of coefficient vectors to be combined.
method	Method for combining vectors of coefficients. The default uses the geometric median.

## Value

A vector of resulting coefficients.

pasteLoc

Get coordinates on plot

## Description

Get coordinates on a plot by mouse-clicking.

#### Usage

```
pasteLoc(nb, digits = c(3, 3))
```

## Arguments

nb	Number of positions.
digits	2 integer indicating the number of decimal places (respectively for x and y co- ordinates).

## Value

A list of coordinates. Note that if you don't put the result in a variable, it returns as the command text for generating the list. This can be useful to get coordinates by mouse-clicking once, but then using the code for convenience and reproducibility.

pcor

## Examples

```
## Not run:
plot(runif(20, max = 5000))
# note the negative number for the rounding of $y
coord <- pasteLoc(3, digits = c(2, -1))
text(coord, c("a", "b", "c"))
## End(Not run)
```

pcor

# Partial correlation

## Description

Partial correlation between x and y, after having adjusted both for z.

# Usage

pcor(x, y, z, alpha = 0.05)

## Arguments

х	A numeric vector.
У	A numeric vector.
z	A data frame, which can contain characters or factors.
alpha	Type-I error for the confidence interval (CI). Default is 0.05, corresponding to a 95% CI.

# Value

The partial correlation, and the lower and upper bounds of its CI.

```
pcor(iris[[1]], iris[[2]], iris[-(1:2)])
```

plot.big\_sp\_list Plot method

## Description

Plot method for class big\_sp\_list.

#### Usage

```
## S3 method for class 'big_sp_list'
plot(x, coeff = 1, ...)
```

#### Arguments

х	An object of class big_sp_list.
coeff	Relative size of text. Default is 1.
	Not used.

#### Value

A ggplot2 object. You can plot it using the print method. You can modify it as you wish by adding layers. You might want to read this chapter to get more familiar with the package ggplot2.

plot.big\_SVD Plot method

## Description

Plot method for class big\_SVD.

## Usage

```
## S3 method for class 'big_SVD'
plot(
    x,
    type = c("screeplot", "scores", "loadings"),
    nval = length(x$d),
    scores = c(1, 2),
    loadings = 1,
    ncol = NULL,
    coeff = 1,
    viridis = TRUE,
    cols = 2,
    ...
)
```

## plot.big\_SVD

#### Arguments

х	An object of class big_SVD.
type	Either
	<ul> <li>"screeplot": plot of decreasing singular values (the default).</li> <li>"scores": plot of the scores associated with 2 Principal Components.</li> <li>"loadings": plot of loadings associated with 1 Principal Component.</li> </ul>
nval	Number of singular values to plot. Default plots all computed.
scores	Vector of indices of the two PCs to plot. Default plots the first two PCs. If providing more than two, it produces many plots.
loadings	Indices of PC loadings to plot. Default plots the first vector of loadings.
ncol	If multiple vector of loadings are to be plotted, this defines the number of columns of the resulting multiplot.
coeff	Relative size of text. Default is 1.
viridis	Deprecated argument.
cols	Deprecated. Use ncol instead.
	Not used.

#### Value

A ggplot2 object. You can plot it using the print method. You can modify it as you wish by adding layers. You might want to read this chapter to get more familiar with the package ggplot2.

#### See Also

big\_SVD, big\_randomSVD and asPlotlyText.

```
set.seed(1)
X <- big_attachExtdata()</pre>
svd <- big_SVD(X, big_scale(), k = 10)</pre>
# screeplots
plot(svd) # 3 PCs seems "significant"
plot(svd, coeff = 1.5) # larger font for papers
# scores plot
plot(svd, type = "scores") # first 2 PCs
plot(svd, type = "scores", scores = c(1, 3))
plot(svd, type = "scores", scores = 1:4, ncol = 2, coeff = 0.7)
## add color (recall that this return a `ggplot2` object)
class(obj <- plot(svd, type = "scores"))</pre>
pop <- rep(c("POP1", "POP2", "POP3"), c(143, 167, 207))</pre>
library(ggplot2)
print(obj2 <- obj + aes(color = pop) + labs(color = "Population"))</pre>
## change the place of the legend
```

#### plot.mhtest

```
print(obj3 <- obj2 + theme(legend.position = c(0.82, 0.17)))
## change the title and the labels of the axes
obj3 + ggtitle("Yet another title") + xlab("with an other 'x' label")
# loadings
plot(svd, type = "loadings", loadings = 2)
## all loadings
plot(svd, type = "loadings", loadings = 1:2, coeff = 0.7, ncol = 1)
# Percentage of variance explained by the PCs
# See https://github.com/privefl/bigstatsr/issues/83
# dynamic plots, require the package **plotly**
## Not run: plotly::ggplotly(obj3)</pre>
```

plot.mhtest Plot method

## Description

Plot method for class mhtest.

#### Usage

```
## S3 method for class 'mhtest'
plot(x, type = c("hist", "Manhattan", "Q-Q", "Volcano"), coeff = 1, ...)
```

#### Arguments

х	An object of class mhtest.
type	Either.
	• "hist": histogram of p-values (the default).
	• "Manhattan": plot of the negative logarithm (in base 10) of p-values.
	• "Q-Q": Q-Q plot.
	• "Volcaco": plot of the negative logarithm of p-values against the estimation of coefficients (e.g. betas in linear regression)
coeff	Relative size of text. Default is 1.
	Not used.

## Value

A ggplot2 object. You can plot it using the print method. You can modify it as you wish by adding layers. You might want to read this chapter to get more familiar with the package ggplot2.

## See Also

big\_univLinReg, big\_univLogReg, plot.big\_SVD and asPlotlyText.

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## predict.big\_sp

set.seed(1)

## Examples

```
X <- big_attachExtdata()
y <- rnorm(nrow(X))
test <- big_univLinReg(X, y)
plot(test)
plot(test, type = "Volcano")
plot(test, type = "Q-Q")
plot(test, type = "Manhattan")
plot(test, type = "Manhattan") + ggplot2::ggtitle(NULL)</pre>
```

predict.big\_sp Predict method

## Description

Predict method for class big\_sp.

## Usage

```
## S3 method for class 'big_sp'
predict(object, X, ind.row, ind.col, covar.row = NULL, ncores = 1, ...)
```

#### Arguments

object	Object of class big_sp.
Х	An object of class FBM.
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
covar.row	Matrix of covariables to be added in each model to correct for confounders (e.g. the scores of PCA), corresponding to ind.row. Default is NULL and corresponds to only adding an intercept to each model. You can use covar_from_df() to convert from a data frame.
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.
	Not used.

## Value

A vector of scores, corresponding to ind.row.

#### See Also

big\_spLinReg and big\_spLogReg.

predict.big\_sp\_list Predict method

# Description

Predict method for class big\_sp\_list.

# Usage

```
## S3 method for class 'big_sp_list'
predict(
    object,
    X,
    ind.row = rows_along(X),
    ind.col = attr(object, "ind.col"),
    covar.row = NULL,
    proba = (attr(object, "family") == "binomial"),
    base.row = NULL,
    ncores = 1,
    ...
)
```

# Arguments

object	Object of class big_sp_list.
Х	An object of class FBM.
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
covar.row	Matrix of covariables to be added in each model to correct for confounders (e.g. the scores of PCA), corresponding to ind.row. Default is NULL and corresponds to only adding an intercept to each model. You can use covar_from_df() to convert from a data frame.
proba	Whether to return probabilities?
base.row	Vector of base predictions, corresponding to ind.row.
ncores	Number of cores used. Default doesn't use parallelism. You may use nb_cores.
	Not used.

## Value

A vector of scores, corresponding to ind.row.

## See Also

big\_spLinReg and big\_spLogReg.

## Description

Get the scores of PCA associated with an svd decomposition (class big\_SVD).

#### Usage

```
## S3 method for class 'big_SVD'
predict(
   object,
   X = NULL,
   ind.row = rows_along(X),
   ind.col = cols_along(X),
   block.size = block_size(nrow(X)),
   ...
)
```

## Arguments

object	A list returned by big_SVD or big_randomSVD.
Х	An object of class FBM.
ind.row	An optional vector of the row indices that are used. If not specified, all rows are used. <b>Don't use negative indices.</b>
ind.col	An optional vector of the column indices that are used. If not specified, all columns are used. <b>Don't use negative indices.</b>
block.size	Maximum number of columns read at once. Default uses block_size.
	Not used.

# Value

A matrix of size  $n \times K$  where n is the number of samples corresponding to indices in ind.row and K the number of PCs computed in object. If X is not specified, this just returns the scores of the training set of object.

#### See Also

predict big\_SVD big\_randomSVD

```
set.seed(1)
```

```
X <- big_attachExtdata()
n <- nrow(X)</pre>
```

```
# Using only half of the data
ind <- sort(sample(n, n/2))</pre>
test <- big_SVD(X, fun.scaling = big_scale(), ind.row = ind)</pre>
str(test)
plot(test$u)
pca <- prcomp(X[ind, ], center = TRUE, scale. = TRUE)</pre>
# same scaling
all.equal(test$center, pca$center)
all.equal(test$scale, pca$scale)
# scores and loadings are the same or opposite
# except for last eigenvalue which is equal to 0
# due to centering of columns
scores <- test$u %*% diag(test$d)</pre>
class(test)
scores2 <- predict(test) # use this function to predict scores</pre>
all.equal(scores, scores2)
dim(scores)
dim(pca$x)
tail(pca$sdev)
plot(scores2, pca$x[, 1:ncol(scores2)])
plot(test$v[1:100, ], pca$rotation[1:100, 1:ncol(scores2)])
# projecting on new data
X2 <- sweep(sweep(X[-ind, ], 2, test$center, '-'), 2, test$scale, '/')</pre>
scores.test <- X2 %*% test$v</pre>
ind2 <- setdiff(rows_along(X), ind)</pre>
scores.test2 <- predict(test, X, ind.row = ind2) # use this</pre>
all.equal(scores.test, scores.test2)
scores.test3 <- predict(pca, X[-ind, ])</pre>
plot(scores.test2, scores.test3[, 1:ncol(scores.test2)])
```

predict.mhtest Predict method

## Description

Predict method for class mhtest.

## Usage

```
## S3 method for class 'mhtest'
predict(object, scores = object$score, log10 = TRUE, ...)
```

## $sub\_bk$

## Arguments

object	An object of class mhtest from you get the probability function with possibly pre-transformation of scores.
scores	Raw scores (before transformation) that you want to transform to p-values.
log10	Are p-values returned on the log10 scale? Default is TRUE.
	Not used.

## Value

Vector of log10(p-values) associated with scores and object.

# See Also

big\_univLinReg and big\_univLogReg.

sub\_bk

Replace extension '.bk'

# Description

Replace extension '.bk'

## Usage

```
sub_bk(path, replacement = "", stop_if_not_ext = TRUE)
```

## Arguments

path	String with extension '.bk'.
replacement	Replacement of '.bk'. Default replaces by nothing.
stop_if_not_ext	
	If replacement != "", whether to error if replacement is not an extension (i.e starting with a dot).

## Value

String with extension '.bk' replaced by replacement.

```
path <- "toto.bk"
sub_bk(path)
sub_bk(path, ".rds")</pre>
```

summary.big\_sp\_list Summary method

## Description

Summary method for class big\_sp\_list.

## Usage

```
## S3 method for class 'big_sp_list'
summary(object, best.only = FALSE, sort = FALSE, ...)
```

## Arguments

object	An object of class big_sp_list.
best.only	Whether to return only one row corresponding to the best model? The best model is the one smallest \$validation_loss.
sort	Whether to sort by \$validation_loss. Default is FALSE.
	Not used.

## Value

A tibble with, for each \$alpha, a mean \$validation\_loss, a mean vector of coefficients \$beta, the corresponding number of non-zero coefficients \$nb\_var, and the reasons of method completion \$message.

|--|

## Description

Theme ggplot2 used by this package.

#### Usage

```
theme_bigstatsr(size.rel = 1)
```

#### Arguments

size.rel Relative size. Default is 1.

```
library(ggplot2)
(p <- ggplot(mapping = aes(x = 1:10, y = 1:10)) + geom_point())
p + theme_bw()
p + theme_bigstatsr()</pre>
```

without\_downcast\_warning

Temporarily disable downcast warning

## Description

Temporarily disable downcast warning

## Usage

without\_downcast\_warning(expr)

# Arguments

expr The expression to evaluate without downcast warning.

## Value

The result of the evaluated expression.

```
without_downcast_warning(FBM(10, 10, type = "integer", init = 1.5))
```

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