Package ‘rsvd’

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Title Randomized Singular Value Decomposition
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Description Low-rank matrix decompositions are fundamental tools and widely used for data analysis, dimension reduction, and data compression. Classically, highly accurate deterministic matrix algorithms are used for this task. However, the emergence of large-scale data has severely challenged our computational ability to analyze big data. The concept of randomness has been demonstrated as an effective strategy to quickly produce approximate answers to familiar problems such as the singular value decomposition (SVD). The rsvd package provides several randomized matrix algorithms such as the randomized singular value decomposition (rsvd), randomized principal component analysis (rpca), randomized robust principal component analysis (rrpca), randomized interpolative decomposition (rid), and the randomized CUR decomposition (rcur). In addition several plot functions are provided.
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**Description**

Subsampled MNIST database of handwritten digits. This smaller dataset has 3000 samples for each of the digits corresponding to the class labels 0,1,2,3. Each 28x28 image patch is stored as a flattened row vector.

**Usage**

```r
data('digits')
```

**Format**

An object of class `rsvd`.

**Source**

`mnist`

**References**

Examples

```r
## Not run:
library('rsvd')
data('digits')

# Display first digit
digit <- matrix(digits[,1], nrow = 28, ncol = 28)
image(digit[,28:1], col = gray(255:0/255))

## End(Not run)
```

---

**ggbiplot**

_Biplot for _rpca _using _ggplot._

**Description**

Creates a pretty biplot which is showing the individual factor map overlayed by the variables factor map, i.e., plotting both the principal component scores and directions.

**Usage**

```r
ggbiplot(
  rpcaObj,
  pcs = c(1, 2),
  loadings = TRUE,
  groups = NULL,
  alpha = 0.6,
  ellipse = TRUE,
  alpha.ellipse = 0.2,
  var_labels = TRUE,
  var_labels.names = NULL,
  ind_labels = TRUE,
  ind_labels.names = NULL
)
```

**Arguments**

- `rpcaObj`:
  - Object returned by the _rpca_ function.

- `pcs`:
  - Array_like.
  - An array with two values indicating the two PCs which should be used for plotting. By default the first two PCs are used, e.g., _c(1, 2)._ 

- `loadings`:
  - _Bool_ (_TRUE, _FALSE_), optional.
  - If _TRUE_, the eigenvectors are unit scaled by the square root of the eigenvalues _W = W * diag(sqrt(eigvals))._

- `groups`:
  - _Factor_, optional.
  - _Factor_ indicating groups.
alpha
Scalar, optional.
Alpha transparency for scatter plot.

ellipse
Bool (TRUE, FALSE), optional.
Draw a 1sd data ellipse for each group, if TRUE.

alpha.ellipse
Scalar, optional.
Alpha transparency for ellipse.

var_labels
Bool (TRUE, FALSE), optional.
Plot variable names, if TRUE.

var_labels.names
Array_like, optional.
User specific labels for the individuals.

ind_labels
Bool (TRUE, FALSE), optional.
Plot data point names, if TRUE.

ind_labels.names
Array_like, optional.
User specific labels for data points.

Author(s)
N. Benjamin Erichson, <erichson@berkeley.edu>

See Also
rpca, ggplot

Examples
#See ?rpca

ggcorplot

Description
Creates a pretty plot which is showing the correlation of the original variable with the principal component (PCs).

Usage
ggcorplot(
rpcaObj,
pcs = c(1, 2),
loadings = TRUE,
var_labels = FALSE,
var_labels.names = NULL,
alpha = 1,
top.n = NULL
)

Examples
#See ?rpca
Arguments

rpcaObj  Object returned by the rpca function.
pcs  Array_like.
An array with two values indicating the two PCs which should be used for plotting. By default the first two PCs are used, e.g., c(1, 2).
loadings  Bool (TRUE, FALSE), optional.
If TRUE, the eigenvectors are unit scaled by the square root of the eigenvalues
\[ W = W \times \text{diag} (\sqrt{\text{eigvals}}) \].
var_labels  Bool (TRUE, FALSE), optional.
Plot variable names, if TRUE.
var_labels.names  Array_like, optional.
User specific labels for the variables
alpha  Scalar, optional.
Alpha transparency of the arrows.
top.n  Scalar, optional.
Number of (most influential) variables to label with small circles.

Author(s)

N. Benjamin Erichson, <erichson@berkeley.edu>

See Also

rpca, ggplot

Examples

#

---

**ggindplot**  *Individual factor map for rpca using ggplot.*

Description

Creates a pretty plot which is showing the individual factor map, i.e, plotting the principal component scores.
Usage

```r
ggindplot(
  rpcaObj,
  pcs = c(1, 2),
  groups = NULL,
  alpha = 0.6,
  ellipse = TRUE,
  alpha.ellipse = 0.2,
  ind_labels = TRUE,
  ind_labels.names = NULL
)
```

Arguments

- **rpcaObj**: Object returned by the `rpca` function.
- **pcs**: Array_like. An array with two values indicating the two PCs which should be used for plotting. By default the first two PCs are used, e.g., c(1, 2).
- **groups**: Factor, optional. Factor indicating groups.
- **alpha**: Scalar, optional. Alpha transparency for scatter plot.
- **ellipse**: Bool (`TRUE`, `FALSE`), optional. Draw a 1sd data ellipse for each group, if `TRUE`.
- **alpha.ellipse**: Scalar, optional. Alpha transparency for ellipse.
- **ind_labels**: Bool (`TRUE`, `FALSE`), optional. Plot names for each individual point, if `TRUE`.
- **ind_labels.names**: Array_like, optional. User specific labels for the individual points.

Author(s)

N. Benjamin Erichson, <erichson@berkeley.edu>

See Also

- `rpca`, `ggplot`

Examples

```r
#See ?rpca
```
ggscreeplot

Pretty Screeplot

Description

Creates a pretty screeplot using `ggplot`. By default the explained variance is plotted against the number of the principal component. Alternatively the explained variance ratio, the cumulative explained variance ratio, or the eigenvalues can be plotted.

Usage

```r
ggscreeplot(rpcaObj, type = c("var", "ratio", "cum", "eigenvals"))
```

Arguments

- `rpcaObj`: Object returned by the `rpca` function.
- `type`: String `c('var', 'ratio', 'cum', 'eigenvals')`, optional.

Author(s)

N. Benjamin Erichson, <erichson@berkeley.edu>

See Also

- `rpca`
- `ggplot`

Examples

```r
#
```

plot.rpga

Screeplot

Description

Creates a screeplot, variables and individual factor maps to summarize the results of the `rpca` function.

Usage

```r
## S3 method for class 'rpca'
plot(x, ...)
```
rcur

Arguments

A   array_like; numeric \((m, n)\) input matrix (or data frame).
If the data contain \(N\) As na.omit is applied.

k   integer; target rank of the low-rank approximation, i.e., the number of columns/rows to be selected. It is required that \(k\) is smaller or equal to \(\min(m, n)\).

p   integer, optional; oversampling parameter (default \(p = 10\)).

q   integer, optional; number of additional power iterations (default \(q = 0\)).

idx_only   bool, optional; if \((T R U E)\), only the index set \(C\).idx and \(R\).idx is returned, but not the matrices \(C\) and \(R\). This is more memory efficient, when dealing with large-scale data.

rand   bool, optional; if \((T R U E)\), a probabilistic strategy is used, otherwise a deterministic algorithm is used.

See Also

ggscreeplot, ggcorplot, ggindplot

Examples

#
Details

Algorithm for computing the CUR matrix decomposition of a rectangular \( (m, n) \) matrix \( A \), with target rank \( k << \min(m, n) \). The input matrix is factored as

\[
A = C \ast U \ast R
\]

using the \texttt{rid} decomposition. The factor matrix \( C \) is formed using actual columns of \( A \), also called the partial column skeleton. The factor matrix \( R \) is formed using actual rows of \( A \), also called the partial row skeleton.

If \( rand = TRUE \) a probabilistic strategy is used to compute the decomposition, otherwise a deterministic algorithm is used.

Value

\texttt{rcur} returns a list with class \texttt{id} containing the following components:

- \texttt{C} array_like;
  
  column subset \( C = A[C.idx] \); \((m, k)\) dimensional array.

- \texttt{R} Array_like;
  
  row subset \( R = A[R.idx,:] \); \((k, n)\) dimensional array.

- \texttt{U} array_like;
  
  connector matrix; \((k, k)\) dimensional array.

- \texttt{C.idx} array_like;
  
  index set of the \( k \) selected columns used to form \( C \).

- \texttt{R.idx} array_like;
  
  index set of the \( k \) selected rows used to form \( R \).

- \texttt{C.scores} array_like;
  
  scores of the selected columns.

- \texttt{R.scores} array_like;
  
  scores of the selected rows.

Author(s)

N. Benjamin Erichson, <erichson@berkeley.edu>

References


See Also

\texttt{rid}
Examples

```r
## Not run:
# Load test image
data('tiger')

# Compute (column) randomized interpolative decomposition
# Note that the image needs to be transposed for correct plotting
out <- rcur(tiger, k = 150)

# Reconstruct image
tiger.re <- out$C %*% out$U %*% out$R

# Compute relative error
print(norm(tiger-tiger.re, 'F') / norm(tiger, 'F'))

# Plot approximated image
image(tiger.re, col = gray((0:255)/255))
## End(Not run)
```

### rid

**Randomized interpolative decomposition (ID).**

**Description**

Randomized interpolative decomposition.

**Usage**

```r
rid(A, k = NULL, mode = "column", p = 10, q = 0, idx_only = FALSE, rand = TRUE)
```

**Arguments**

- `A` : array_like; numeric \((m, n)\) input matrix (or data frame).
  If the data contain \(N\)As na.omit is applied.
- `k` : integer, optional; number of rows/columns to be selected. It is required that \(k\) is smaller or equal to \(\min(m, n)\).
- `mode` : string c('column', 'row'), optional; columns or rows ID.
- `p` : integer, optional; oversampling parameter (default \(p = 10\)).
- `q` : integer, optional; number of additional power iterations (default \(q = 0\)).
**idx_only**  bool, optional:
    if (TRUE), the index set idx is returned, but not the matrix C or R. This is more memory efficient, when dealing with large-scale data.

**rand**  bool, optional:
    if (TRUE), a probabilistic strategy is used, otherwise a deterministic algorithm is used.

**Details**

Algorithm for computing the ID of a rectangular $(m, n)$ matrix $A$, with target rank $k << \min(m, n)$. The input matrix is factored as

$$A = C \ast Z$$

using the column pivoted QR decomposition. The factor matrix $C$ is formed as a subset of columns of $A$, also called the partial column skeleton. If mode='column', then the input matrix is factored as

$$A = Z \ast R$$

using the row pivoted QR decomposition. The factor matrix $R$ is now formed as a subset of rows of $A$, also called the partial row skeleton. The factor matrix $Z$ contains a $(k, k)$ identity matrix as a submatrix, and is well-conditioned.

If rand != TRUE a probabilistic strategy is used to compute the decomposition, otherwise a deterministic algorithm is used.

**Value**

`rid` returns a list containing the following components:

- **C** array_like;
  column subset $C = A[\text{idx}]$, if mode='column'; array with dimensions $(m, k)$.

- **R** array_like;
  row subset $R = A[\text{idx}]$, if mode='row'; array with dimensions $(k, n)$.

- **Z** array_like;
  well conditioned matrix; Depending on the selected mode, this is an array with dimensions $(k, n)$ or $(m, k)$.

- **idx** array_like;
  index set of the $k$ selected columns or rows used to form $C$ or $R$.

- **pivot** array_like;
  information on the pivoting strategy used during the decomposition.

- **scores** array_like;
  scores of the columns or rows of the input matrix $A$.

- **scores.idx** array_like;
  scores of the $k$ selected columns or rows in $C$ or $R$. 

Description

Fast computation of the principal components analysis using the randomized singular value decomposition.
Usage

```r
rpca(
  A,
  k = NULL,
  center = TRUE,
  scale = TRUE,
  retx = TRUE,
  p = 10,
  q = 2,
  rand = TRUE
)
```

Arguments

- **A** array_like; a numeric $(m,n)$ input matrix (or data frame) to be analyzed. If the data contain NAs na.omit is applied.
- **k** integer; number of dominant principle components to be computed. It is required that $k$ is smaller or equal to $\min(m,n)$, but it is recommended that $k << \min(m,n)$.
- **center** bool, optional; logical value which indicates whether the variables should be shifted to be zero centered (TRUE by default).
- **scale** bool, optional; logical value which indicates whether the variables should be scaled to have unit variance (TRUE by default).
- **retx** bool, optional; logical value indicating whether the rotated variables / scores should be returned (TRUE by default).
- **p** integer, optional; oversampling parameter for rsvd (default $p = 10$), see rsvd.
- **q** integer, optional; number of additional power iterations for rsvd (default $q = 1$), see rsvd.
- **rand** bool, optional; if (TRUE), the rsvd routine is used, otherwise svd is used.

Details

Principal component analysis is an important linear dimension reduction technique.

Randomized PCA is computed via the randomized SVD algorithm (rsvd). The computational gain is substantial, if the desired number of principal components is relatively small, i.e. $k << \min(m,n)$.

The print and summary method can be used to present the results in a nice format. A scree plot can be produced with ggscreeplot. The individuals factor map can be produced with gginfmaplot, and a correlation plot with ggcorplot.
The predict function can be used to compute the scores of new observations. The data will automatically be centered (and scaled if requested). This is not fully supported for complex input matrices.

Value

rpca returns a list with class rpca containing the following components:

- **rotation** array_like;
  the rotation (eigenvectors); \((n, k)\) dimensional array.

- **eigvals** array_like;
  eigenvalues; \(k\) dimensional vector.

- **sdev** array_like;
  standard deviations of the principal components; \(k\) dimensional vector.

- **x** array_like;
  the scores / rotated data; \((m, k)\) dimensional array.

- **center, scale** array_like;
  the centering and scaling used.

Note

The principal components are not unique and only defined up to sign (a constant of modulus one in the complex case) and so may differ between different PCA implementations.

Similar to prcomp the variances are computed with the usual divisor \(N - 1\).

Author(s)

N. Benjamin Erichson, <erichson@berkeley.edu>

References


See Also

ggscreepplot, ggindplot, ggcorplot, plot.rpca, predict.rsvd

Examples

```r
library('rsvd')
#
# Load Edgar Anderson's Iris Data
#
```
```r
data('iris')

# # log transform #
log.iris <- log( iris[, 1:4] )
iris.species <- iris[, 5]

# # Perform rPCA and compute only the first two PCs #
iris.rpca <- rpca(log.iris, k=2)
summary(iris.rpca) # Summary
print(iris.rpca) # Prints the rotations

# # Use rPCA to compute all PCs, similar to \code{\link[prcomp]} #
iris.rpca <- rpca(log.iris)
summary(iris.rpca) # Summary
print(iris.rpca) # Prints the rotations
plot(iris.rpca) # Produce screeplot, variable and individuals factor maps.
```

---

**rqb**  
*Randomized QB Decomposition (rqb).*

**Description**

Compute the near-optimal QB decomposition of a rectangular matrix.

**Usage**

```r
rqb(A, k = NULL, p = 10, q = 2, sdist = "normal", rand = TRUE)
```

**Arguments**

- **A**  
  array_like;  
  real/complex \((m, n)\) input matrix (or data frame).
- **k**  
  integer, optional;  
  target rank of the low-rank decomposition. It should satisfy \(k << \min(m, n)\).
- **p**  
  integer, optional;  
  oversampling parameter (default \(p = 10\)).
- **q**  
  integer, optional;  
  number of power iterations (default \(q = 2\)).
sdist is a string that specifies the sampling distribution:
- `'unif'`: Uniform `[-1,1]`.
- `'normal'` (default): Normal `~N(0,1)`.
- `'rademacher'`: Rademacher random variates.

rand is a boolean that, if `TRUE`, uses a probabilistic strategy; otherwise a deterministic algorithm is used.

**Details**

The randomized QB decomposition factors a rectangular \((m,n)\) matrix \(A\) as \(A = Q \ast B\). \(Q\) is an \((m,k)\) matrix with orthogonal columns, and \(B\) a \((k,n)\) matrix. The target rank is assumed to be \(k << \min(m,n)\).

\(p\) is an oversampling parameter to improve the approximation. A value between 5 and 10 is recommended, and \(p = 10\) is set by default.

The parameter \(q\) specifies the number of power (subspace) iterations to reduce the approximation error. This is recommended if the singular values decay slowly. In practice 1 or 2 iterations achieve good results, however, computing power iterations increases the computational time. The number of power iterations is set to \(q = 2\) by default.

**Value**

\(rqb\) returns a list containing the following components:

- \(Q\) array_like:
  - matrix with orthogonal columns; \((m,k)\) dimensional array.

- \(B\) array_like:
  - smaller matrix; \((k,n)\) dimensional array.

**Author(s)**

N. Benjamin Erichson, <erichson@berkeley.edu>

**References**


**See Also**

- `svd`
rrpca

Randomized robust principal component analysis (rrpca).

Description

Robust principal components analysis separates a matrix into a low-rank plus sparse component.

Usage

```r
rrpca(
  A,
  lambda = NULL,
  maxiter = 50,
  tol = 1e-05,
  p = 10,
  q = 2,
  trace = FALSE,
  rand = TRUE
)
```

Arguments

- **A**: array_like; a real $(m, n)$ input matrix (or data frame) to be decomposed. na.omit is applied, if the data contain NAs.
- **lambda**: scalar, optional; tuning parameter (default $\lambda = \max(m, n)^{-0.5}$).
- **maxiter**: integer, optional; maximum number of iterations (default $\text{maxiter} = 50$).
- **tol**: scalar, optional; precision parameter (default $\text{tol} = 1.0 \times 10^{-5}$).
- **p**: integer, optional; oversampling parameter for rsvd (default $p = 10$), see rsvd.
- **q**: integer, optional; number of additional power iterations for rsvd (default $q = 2$), see rsvd.
- **trace**: bool, optional; print progress.
- **rand**: bool, optional; if (TRUE), the rsvd routine is used, otherwise svd is used.

Details

Robust principal component analysis (RPCA) is a method for the robust separation of a rectangular $(m, n)$ matrix $A$ into a low-rank component $L$ and a sparse component $S$: 
\[ A = L + S \]

To decompose the matrix, we use the inexact augmented Lagrange multiplier method (IALM). The algorithm can be used in combination with either the randomized or deterministic SVD.

**Value**

`rrpca` returns a list containing the following components:

- **L** array_like; low-rank component; \((m, n)\) dimensional array.
- **S** array_like; sparse component; \((m, n)\) dimensional array.

**Author(s)**

N. Benjamin Erichson, <erichson@berkeley.edu>

**References**


**Examples**

```r
library('rsvd')

# Create toy video
# background frame
xy <- seq(-50, 50, length.out=100)
mgrid <- list(x=outer(xy*0,xy,FUN="+"), y=outer(xy,xy*0,FUN="+"))
bg <- 0.1*exp(sin(-mgrid$x**2-mgrid$y**2))
toyVideo <- matrix(rep(c(bg), 100), 100*100, 100)

# add moving object
for(i in 1:90) {
  mobject <- matrix(0, 100, 100)
  mobject[i:(10+i), 45:55] <- 0.2
  toyVideo[,i] = toyVideo[,i] + c(mobject)
}

# Foreground/Background separation
out <- rrpca(toyVideo, trace=TRUE)

# Display results of the seperation for the 10th frame
```
rsvd

Randomized Singular Value Decomposition (rsvd).

Description

The randomized SVD computes the near-optimal low-rank approximation of a rectangular matrix using a fast probabilistic algorithm.

Usage

rsvd(A, k = NULL, nu = NULL, nv = NULL, p = 10, q = 2, sdist = "normal")

Arguments

A
array_like;
a real/complex \((m, n)\) input matrix (or data frame) to be decomposed.

k
integer;
specifies the target rank of the low-rank decomposition. \(k\) should satisfy \(k << \min(m, n)\).

nu
integer, optional;
number of left singular vectors to be returned. \(nu\) must be between 0 and \(k\).

nv
integer, optional;
number of right singular vectors to be returned. \(nv\) must be between 0 and \(k\).

p
integer, optional;
oversampling parameter (by default \(p = 10\)).

q
integer, optional;
number of additional power iterations (by default \(q = 2\)).

sdist
string \(c(\text{'unif'}, \text{'normal'}, \text{'rademacher'})\), optional;
specifies the sampling distribution of the random test matrix:
\(\text{'unif'}\) : Uniform \([-1,1]\).
\(\text{'normal'}\) (default) : Normal \(\sim \text{N}(0,1)\).
\(\text{'rademacher'}\) : Rademacher random variates.

Details

The singular value decomposition (SVD) plays an important role in data analysis, and scientific computing. Given a rectangular \((m, n)\) matrix \(A\), and a target rank \(k << \min(m, n)\), the SVD factors the input matrix \(A\) as
\[ A = U_k \text{diag}(d_k)V_k^\top \]

The \( k \) left singular vectors are the columns of the real or complex unitary matrix \( U \). The \( k \) right singular vectors are the columns of the real or complex unitary matrix \( V \). The \( k \) dominant singular values are the entries of \( d \), and non-negative and real numbers.

\( p \) is an oversampling parameter to improve the approximation. A value of at least 10 is recommended, and \( p = 10 \) is set by default.

The parameter \( q \) specifies the number of power (subspace) iterations to reduce the approximation error. The power scheme is recommended, if the singular values decay slowly. In practice, 2 or 3 iterations achieve good results, however, computing power iterations increases the computational costs. The power scheme is set to \( q = 2 \) by default.

If \( k > (\min(n,m)/4) \), a deterministic partial or truncated \( \text{svd} \) algorithm might be faster.

**Value**

\( \text{rsvd} \) returns a list containing the following three components:

- \( d \) array_like;
  singular values; vector of length \( k \).
- \( u \) array_like;
  left singular vectors; \((m,k)\) or \((m,nu)\) dimensional array.
- \( v \) array_like;
  right singular vectors; \((n,k)\) or \((n,nv)\) dimensional array.

**Note**

The singular vectors are not unique and only defined up to sign (a constant of modulus one in the complex case). If a left singular vector has its sign changed, changing the sign of the corresponding right vector gives an equivalent decomposition.

**Author(s)**

N. Benjamin Erichson, <erichson@berkeley.edu>

**References**


**See Also**

- \( \text{svd} \), \( \text{rpca} \)
Examples

```r
library('rsvd')

# Create a n x n Hilbert matrix of order n,
# with entries H[i,j] = 1 / (i + j + 1).
hilbert <- function(n) { i <- 1:n; 1 / outer(i - 1, i, "+") }
H <- hilbert(n=50)

# Low-rank (k=10) matrix approximation using rsvd
k=10
s <- rsvd(H, k=k)
Hre <- s$u %*% diag(s$d) %*% t(s$v) # matrix approximation
print(100 * norm( H - Hre, 'F') / norm( H,'F')) # percentage error

# Compare to truncated base svd
s <- svd(H)
Hre <- s$u[,1:k] %*% diag(s$d[,1:k]) %*% t(s$v[,1:k]) # matrix approximation
print(100 * norm( H - Hre, 'F') / norm( H,'F')) # percentage error
```

---

tiger

**Tiger**

Description

1600x1200 grayscaled (8 bit [0-255]/255) image.

Usage

```r
data('tiger')
```

Format

An object of class `rsvd`.

Source

Wikimedia

References

S. Taheri (2006). "Panthera tigris altaica", (Online image)

Examples

```r
## Not run:
library('rsvd')
data('tiger')

#Display image
```
image(tiger, col = gray((0:255)/255))

## End(Not run)
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