

Package: blox (via r-universe)

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Type Package

Title Block Diagonal Matrix Approximation

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Description Finds the best block diagonal matrix approximation of a symmetric matrix. This can be exploited for divisive hierarchical clustering using singular vectors, named HC-SVD. The method is described in Bauer (202Xa) <[doi:10.48550/arXiv.2308.06820](https://doi.org/10.48550/arXiv.2308.06820)>.

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bd.approx

*Two Block Diagonal Matrix Approximation.***Description**

Finds the best two block diagonal matrix approximation of a similarity matrix according to some distance (linkage) function as described in Bauer (202Xa). Candidate splits are determined by the first sparse eigenvectors (sparse approximations of the first eigenvectors, i.e., vectors with many zero entries) of the similarity matrix.

Usage

```
bd.approx(S, linkage = "average", q = 1, h.power = 2, balance, max.iter)
```

Arguments

S	A scaled $p \times p$ similarity matrix. For example, this may be a correlation matrix.
linkage	The linkage function to be used. This should be one of "average", "ncut", "rcut", "RV" (for RV-coefficient), or "single".
q	Number of sparse eigenvectors to be used. This should be either a numeric value between zero and one to indicate percentages, or "Kaiser" for as many sparse eigenvectors as there are eigenvalues larger or equal to one. For a numerical value between zero and one, the number of sparse eigenvectors is determined as the corresponding share of the total number of eigenvectors. E.g., $q = 1$ (100%) uses all sparse eigenvectors and $q = 0.5$ (50%) will use half of all sparse eigenvectors. For $q = 1$, identification is best (see Bauer (202Xa) for details).
h.power	h -th Hadamard power of S . This should be a positive integer and increases robustness of the method, as described in Bauer (202Xa).
balance	Minimum proportion of the smaller block when splitting into two. Must be a numeric value in $(0, 0.5]$. For example, $balance = 0.5$ enforces an exact 50:50 split, while $balance = 0.2$ allows splits as unbalanced as 20:80, with more balanced splits such as 30:70 or 40:60 also permitted. If an exact split is not possible (e.g., $balance = 0.5$ when $p = 9$), the closest integer partition is used (e.g., 4 and 5 per block).
max.iter	How many iterations should be performed for computing the sparse eigenvectors. Default is 500.

Details

The sparse eigenvectors are computed using the method of Shen and Huang (2008). The method is implemented by Baglama, Reichel, and Lewis in `ssvd` (`ir1ba`). Here, we use a Rcpp/RcppArmadillo implementation based on `ssvd` with slight modifications to suit our method and for faster performance.

Value

A list with four components:

B	The best two block diagonal matrix approximation.
BD	The best two block diagonal matrix approximation permuted to a block diagonal shape: $BD = PBt(P)$.
P	The permutation matrix P : $BD = PBt(P)$.
clustering	The clustering vector as an integer vector of length p , which gives for each component the number 1 or 2 of the cluster/split to which it belongs.
split	A list containing the two splits.
distance	The approximation error (distance) according to the selected linkage function.

References

Bauer, J.O. (202Xa). Divisive hierarchical clustering using block diagonal matrix approximations. Working paper.

Shen, H. and Huang, J.Z. (2008). Sparse principal component analysis via regularized low rank matrix approximation, J. Multivar. Anal. 99, 1015–1034.

Examples

```
#We give a trivial example for a block diagonal matrix perturbed by
#noise, for adapting clustering objectives of spectral clustering,
#and for balanced clustering.
```

```
### TOY EXAMPLE
```

```
A <- matrix(c(2,1,1,3), 2, 2) # 2x2 block
B <- matrix(c(5,4,4,6), 2, 2) # 2x2 block
```

```
# Create a 5x5 zero matrix and insert blocks at right positions.
M <- matrix(0, 4, 4)
M[1:2, 1:2] <- A
M[3:4, 3:4] <- B
```

```
M.tilde <- M + matrix(rnorm(4^2, 0, 0.2), 4, 4)
```

```
#Construct a similarity matrix with same block structure
S <- cov2cor(t(M.tilde) %*% M.tilde)
bd <- bd.approx(S)
```

```
#Block diagonal approximation:
bd$B
```

```
#We can also permute the block diagonal shape:
S2 <- S[c(1, 3, 2, 4), c(1, 3, 2, 4)]
bd2 <- bd.approx(S2)
```

```

#bd2$B gives us again the block diagonal approximation
bd2$B

#And bd2$BD gives us the block diagonal approximation permuted to
#block diagonal shape
bd2$BD

### ADAPTING CLUSTERING OBJECTIVES

#We will use the USArrests example (see ?hcsvd).
data("USArrests")
USArrests["Maryland", "UrbanPop"] <- 76.6
D <- as.matrix(dist(USArrests))
S <- 1 - D / max(D)

#We compute k = 2 clusters adapting the objective of spectral clustering
#with the ratio cut.
bd.approx(S, linkage = "rcut")

### BALANCED CLUSTERING

#We can also enforce balanced clustering, such as two clusters of equal
#size (50:50). We will do this for the USArrests example from above.
bd.approx(S, linkage = "rcut", balance = 0.5)

```

hc.beta

Compute Revelle's Beta for all worst split-halves using HC-SVD.

Description

Performs HC-SVD to reveal the hierarchical variable structure using average linkage as described in Bauer (202Xa). For a data matrix comprising p items, this means that $p - 1$ splits are identified. The obtained structure aligns with the structure according to the worst split-half reliability and is thus used to compute a hierarchy of all Revelle's beta as described in Bauer (202Xb).

Usage

```
hc.beta(R, splits = NULL, n.splits = NULL, is.corr = TRUE, verbose = TRUE)
```

Arguments

R A correlation matrix of dimension pxp or a data matrix of dimension npx can be provided. If a data matrix is supplied, it must be indicated by setting `is.corr = FALSE`, and the correlation matrix will then be calculated as `cor(X)`.

splits	An object containing the splits identified by HC-SVD. This can either be the result of <code>hcsvd</code> (for all splits) or <code>bd.approx</code> (for a single split). If omitted, <code>hc.beta</code> will internally call <code>hcsvd(R)</code> and compute Revelle's beta for all $p - 1$ splits.
n.splits	Number of splits for which Revelle's beta is computed. If <code>splits</code> is from <code>hcsvd</code> , the default is all $p - 1$ splits. If <code>splits</code> is from <code>bd.approx</code> , only a single split is available and <code>n.splits</code> is set to 1.
is.corr	Is the supplied object a correlation matrix. Default is TRUE and this parameter must be set to FALSE if a data matrix instead of a correlation matrix is supplied.
verbose	Print out progress as $p - 1$ iterations for divisive hierarchical clustering are performed. Default is TRUE.

Details

Supplementary details are in Bauer (202Xb).

Value

A list with `n.splits` components. Each split is a list of four components:

split	The split number.
beta	Revelle's beta for this split.
A	One of the two sub-scales that has been split.
B	One of the two sub-scales that has been split.
beta.alpha	Computes the ratio of Revelle's beta and Cronbach's alpha.

References

Bauer, J.O. (202Xa). *Divisive hierarchical clustering using block diagonal matrix approximations. Working paper.*

Bauer, J.O. (202Xb). *Revelle's beta: The wait is over - we can compute it! Working paper.*

See Also

[bd.approx](#) [hcsvd](#)

Examples

```
#We compute the worst split-half reliabilities on a correlation matrix.
```

```
#Load the correlation matrix Bechtoldt from the psych
#package (see ?Bechtoldt for more information).
if (requireNamespace("psych", quietly = TRUE)) {
  data("Bechtoldt", package = "psych")
}
R <- Bechtoldt
```

```
### RUN HC-SVD FOR HIERARCHICAL VARIABLE CLUSTERING

#Compute HC-SVD (with average linkage).
hcsvd.obj <- hcsvd(R)

#The object of type hclust with corresponding dendrogram can be obtained
#directly from hcsvd(...):
hc.div <- hcsvd.obj$hclust
plot(hc.div, axes = FALSE, ylab = "", main = "Revelle's Beta Splits")

### COMPUTE REVELLE'S BETA FOR ALL IDENTIFIED SPLITS

#Compute Revelle's beta
betas <- hc.beta(R = R)

#Alternatively, you can submit the object obtained from hcsvd(). Thus,
#the hierarchy needs not to be computed again using hcsvd().
betas <- hc.beta(R = R, splits = hcsvd.obj)

#Visualize the splits, e.g., as
splits <- sapply(betas, `[`, "split")
beta.values <- sapply(betas, `[`, "beta")

plot(splits, beta.values,
     type = "b",
     xlab = "Split",
     ylab = "Revelle's Beta",
     main = "Revelle's Beta Across Splits",
     pch = 19)

#Visualize the ratio of Revelle's beta and Cronbach's alpha
beta.alpha <- sapply(betas, `[`, "beta.alpha")
plot(splits, beta.values,
     type = "b",
     xlab = "Split",
     ylab = "Beta/Alpha",
     main = "Ratio of Beta and Alpha Across Splits",
     pch = 19)

### COMPUTE REVELLE'S BETA FOR THE FIRST IDENTIFIED SPLIT

#The first split can be identified using bd.approx()
#This is computationally faster, as only the first split
#is identified
hc.beta(R = R, splits = bd.approx(R))
```

Description

Performs HC-SVD to reveal the hierarchical structure as described in Bauer (202Xa). This divisive approach iteratively splits each cluster into two subclusters. Candidate splits are determined by the first sparse eigenvectors (sparse approximations of the first eigenvectors, i.e., vectors with many zero entries) of the similarity matrix. The selected split is the one that yields the best block-diagonal approximation of the similarity matrix according to a specified linkage function. The procedure continues until each object is assigned to its own cluster.

Usage

```
hcsvd(S, linkage = "average", q = 1, h.power = 2, max.iter, verbose = TRUE)
```

Arguments

S	A scaled $p \times p$ similarity matrix. For example, this may be a correlation matrix.
linkage	The linkage function to be used. This should be one of "average", "single", or "RV" (for RV-coefficient). Note that the RV-coefficient might not yield an ultrametric distance.
q	Number of sparse eigenvectors to be used. This should be either a numeric value between zero and one to indicate percentages, or "Kaiser" for as many sparse eigenvectors as there are eigenvalues larger or equal to one. For a numerical value between zero and one, the number of sparse eigenvectors is determined as the corresponding share of the total number of eigenvectors. E.g., $q = 1$ (100%) uses all sparse eigenvectors and $q = 0.5$ (50%) will use half of all sparse eigenvectors. For $q = 1$, identification is best (see Bauer (202Xa) for details).
h.power	h -th Hadamard power of S . This should be a positive integer and increases robustness of the method, as described in Bauer (202Xa).
max.iter	How many iterations should be performed for computing the sparse eigenvectors. Default is 500.
verbose	Print out progress as $p - 1$ iterations for divisive hierarchical clustering are performed. Default is TRUE.

Details

The sparse loadings are computed using the method proposed by Shen & Huang (2008). The corresponding implementation is written in Rcpp/RcppArmadillo for computational efficiency and is based on the R implementation by Baglama, Reichel, and Lewis in `ssvd` (`irlba`). However, the implementation has been adapted to better align with the scope of the `bdsvd` package which is the base for the `blox` package.

Supplementary details are in [hc.beta](#) and in Bauer (202Xb).

Value

A list with four components:

<code>hclust</code>	The clustering structure identified by HC-SVD as an object of type <code>hclust</code> .
<code>dist.matrix</code>	The ultrametric distance matrix (cophenetic matrix) of the HC-SVD structure as an object of class <code>dist</code> .
<code>u.sim</code>	The ultrametric similarity matrix of S obtained by HC-SVD as an object of class <code>matrix</code> . The ultrametric similarity matrix is calculated as <code>1-dist.matrix</code> .
<code>q.p</code>	A vector of length $p - 1$ containing the ratio q_i/p_i of the q_i sparse eigenvectors used relative to all sparse eigenvectors q_i for the split of each cluster. The ratio is set to NA if the cluster contains only two variables as the search for sparse eigenvectors that reflect this obvious split is not required in this case.

References

- Bauer, J.O. (202Xa). Divisive hierarchical clustering using block diagonal matrix approximations. Working paper.*
- Bauer, J.O. (202Xb). Revelle's beta: The wait is over - we can compute it!. Working paper.*
- Shen, H. and Huang, J.Z. (2008). Sparse principal component analysis via regularized low rank matrix approximation, J. Multivar. Anal. 99, 1015–1034.*

See Also

[bdsvd](#) {`bdsvd`}

Examples

```
#We give one example for variable clustering directly on a correlation matrix,
#and we replicate the USArrest example in Bauer (202Xa) for observation clustering.
#More elaborate code alongside a different example for variable clustering can be
#found in the corresponding supplementary material of that manuscripts.

### VARIABLE CLUSTERING

#Load the correlation matrix Bechtoldt from the psych
#package (see ?Bechtoldt for more information).
if (requireNamespace("psych", quietly = TRUE)) {
  data("Bechtoldt", package = "psych")
}

#Compute HC-SVD (with average linkage).
hcsvd.obj <- hcsvd(Bechtoldt)

#The object of type hclust with corresponding dendrogram can be obtained
#directly from hcsvd(...):
hc.div <- hcsvd.obj$hclust
plot(hc.div, ylab = "")
```

```
#The dendrogram can also be obtained from the ultrametric distance matrix:
plot(hclust(hcsvd.obj$dist.matrix), main = "HC-SVD", sub = "", xlab = "")

### OBSERVATION CLUSTERING

#Correct for the known transcription error
data("USArrests")
USArrests["Maryland", "UrbanPop"] <- 76.6

#The distance matrix is scaled (divided by max(D)) to later allow a
#transformation to a matrix S that fulfills the properties of a similarity
#matrix.
D <- as.matrix(dist(USArrests))
D <- D / max(D)
S <- 1 - D

#Compute HC-SVD (with average linkage).
hcsvd.obj <- hcsvd(S)

#The object of type hclust with corresponding dendrogram can be obtained
#directly from hcsvd(...):
hc.div <- hcsvd.obj$hclust
plot(hc.div, ylab = "")

#The dendrogram can also be obtained from the ultrametric distance matrix:
plot(hclust(hcsvd.obj$dist.matrix), main = "HC-SVD", sub = "", xlab = "")
```

is.ultrametric	<i>Ultrametric Distance Property</i>
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Description

This function checks the ultrametric property of a distance matrix obtained by HC-SVD.

Usage

```
is.ultrametric(hcsvd.obj)
```

Arguments

hcsvd.obj An object of type hcsvd(...)

Value

Returns TRUE if the ultrametric property is fulfilled. Otherwise FALSE.

References

Bauer, J.O. (202Xa). Divisive hierarchical clustering using block diagonal matrix approximations. Working paper.

See Also

[hcsvd](#)

Examples

```
#Load the correlation matrix Bechtoldt from the psych
#package (see ?Bechtoldt for more information).
if (requireNamespace("psych", quietly = TRUE)) {
  data("Bechtoldt", package = "psych")
}

#Compute HC-SVD (with average linkage).
hcsvd.obj <- hcsvd(Bechtoldt)

#Check the ultrametric property
is.ultrametric(hcsvd.obj)
```

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