Package ‘SpatialBSS’

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Description Blind source separation for multivariate spatial data based on simultaneous/joint diagonalization of (robust) local covariance matrices. This package is an implementation of the methods described in Bachoc, Genton, Nordhausen, Ruiz-Gazen and Virta (2020) <doi:10.1093/biomet/asz079>.
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**Description**


**Details**

- **Package:** SpatialBSS
- **Type:** Package
- **Version:** 0.14-0
- **Date:** 2023-07-20
- **License:** GPL (>= 2)

This package provides functions to solve the Blind Source Separation problem for multivariate spatial data. These methods are designed to work with random fields that are observed on irregular locations. Moreover, the random field is assumed to show weak second order stationarity. The main functions of this package are:

- **sbss** This function derives a set of local scatter matrices that are based on spatial kernel functions, where the spatial kernel functions can be chosen. Then this set of local covariance matrices as well as the sample covariance matrix are simultaneously/jointly diagonalized. Local covariance matrices as well as local difference matrices are implemented.
• **sbss_asym**, **sbss_boot** These functions test for white noise components in the estimated latent field estimated by the `sbss` function based on asymptotic results or bootstrap inference principles.

• **snss_sd**, **snss_jd** and **snss_sjd** These functions estimate the latent random field assuming a spatial non-stationary source separation model. This is done by splitting the domain into a number of sub-domains and diagonalizing the corresponding covariance and/or local covariance matrices for each sub-domain.

• **robsbss** Uses robust estimates of local covariance matrices to solve the SBSS problem.

Joint diagonalization is computed with the `frjd` (fast real joint diagonalization) algorithm from the package **JADE**.

The random field can be either a pair of numeric matrices giving the coordinates and field values or an object of class `SpatialPointsDataFrame` or `sf`.

**Author(s)**

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**References**


---

```r
coef.sbss
```

**Description**

Extracts the estimated unmixing matrix of an object of class 'sbss'.

**Usage**

```r
## S3 method for class 'sbss'
coef(object, ...)```
**Arguments**

- `object` object of class 'sbss'. Usually result of `sbss`.
- ... further arguments to be passed to or from methods.

**Value**

Returns the estimated unmixing matrix of an object of class 'sbss' as a numeric matrix.

**See Also**

`sbss`

---

**Description**

Generates synthetic global outliers and contaminates a given p-variate random field

**Usage**

```r
gen_glob_outl(x, alpha = 0.05, h = 10, random_sign = FALSE)
```

**Arguments**

- `x` a numeric matrix of dimension `c(n, p)` where the `p` columns correspond to the entries of the random field and the `n` rows are the observations.
- `alpha` a numerical value between 0 and 1 giving the proportion of observations to contaminate.
- `h` a numerical constant to determine how large the contaminated outliers are, see details.
- `random_sign` logical. If TRUE, the sign of each component of the outlier is randomly selected. Default is FALSE. See more in details.

**Details**

`gen_glob_outl` generates outliers for a given field by selecting randomly round(alpha * n) observations $x_i$ to be the outliers and contaminating them by setting $x_i^{out} = (c')'x_i$, where the elements $c_j$ of vector $c'$ are determined by the parameter `random_sign`. If `random_sign = TRUE`, $c_j$ is either $h$ or $-h$ with $P(c_j = h) = P(c_j = -h) = 0.5$. If `random_sign = FALSE`, $c_j = h$ for all $j = 1,...,p$, $i = 1,...,n$. The parameter `alpha` determines the contamination rate $\alpha$ and the parameter `h` determines the size of the outliers.

**Value**

`gen_glob_outl` returns a `data.frame` containing the contaminated fields as `p` first columns. The column `p + 1` contains a logical indicator whether the observation is outlier or not.
gen_loc_outl

See Also

- gen_loc_outl

Examples

```r
# simulate coordinates
coords <- runif(1000 * 2) * 20
dim(coords) <- c(1000, 2)
coords_df <- as.data.frame(coords)
names(coords_df) <- c("x", "y")
# simulate random field
if (!requireNamespace("gstat", quietly = TRUE)) {
  message("Please install the package gstat to run the example code."
} else {
  library(gstat)
  model_1 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                   model = vgm(psill = 0.025, range = 1, model = 'Exp'), nmax = 20)
  model_2 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                   model = vgm(psill = 0.025, range = 1, kappa = 2, model = 'Mat'),
                   nmax = 20)
  model_3 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                   model = vgm(psill = 0.025, range = 1, model = 'Gau'), nmax = 20)
  field_1 <- predict(model_1, newdata = coords_df, nsim = 1)$sim1
  field_2 <- predict(model_2, newdata = coords_df, nsim = 1)$sim1
  field_3 <- predict(model_3, newdata = coords_df, nsim = 1)$sim1
  field <- cbind(field_1, field_2, field_3)
  # Generate 10 % global outliers to data, with size h=15.
  field_cont <- gen_glob_outl(field, alpha = 0.1, h = 15)
  # Generate 5 % global outliers to data, with size h = 10 and random sign.
  field_cont2 <- gen_glob_outl(field, alpha = 0.05, h = 10, random_sign = TRUE)
}
```

---

Contamination with Local Outliers

Description

Generates synthetic local outliers and contaminates a given p-variate random field by swapping observations based on the first principal component score.

Usage

```r
gen_loc_outl(x, coords, alpha = 0.05,
               neighborhood_type = c("radius", "fixed_n"),
               radius = NULL,
               neighborhood_size = NULL,
               swap_order = c("regular", "reverse", "random"))
```
Arguments

- **x**: a numeric matrix of dimension $c(n, p)$ where the $p$ columns correspond to the entries of the random field and the $n$ rows are the observations.
- **coords**: a numeric matrix or data frame with dimension $c(n, 2)$ containing the coordinates of the observations.
- **alpha**: a numeric value between 0 and 1 determining the proportion of the contaminated observations.
- **neighborhood_type**: a string determining the type of neighborhood. If 'radius', each neighborhood contains all points within the radius determined by the parameter radius. If 'fixed_n', each neighborhood contains a constant number of closest points, where the constant is determined by the parameter neighborhood_size. Default is 'radius'.
- **radius**: a positive numeric value defining the size of the radius when the neighborhood_type is 'radius'. If NULL the radius defaults as $0.01 \times n$.
- **neighborhood_size**: a positive integer defining the number of points in each neighborhood when the neighborhood_type is 'fixed_n'. If NULL the number of points defaults as ceiling($0.01 \times n$).
- **swap_order**: a string to determine which swap order is used. Either 'regular' (default), 'reverse' or 'random'. See details.

Details

gen_loc_outl generates local outliers by swapping the most extreme and the least extreme observations based on the first principal component score under the condition that at most one outliers lies in each neighborhood. For each location $s_i$, the neighborhood $N_i$ is defined based on the parameter neighborhood_type. When neighborhood_type is 'radius', the neighborhood $N_i$ contains all locations $s_j$ for which the Euclidean norm $||s_i - s_j|| < r$, where $r$ is determined by the parameter radius. When neighborhood_type is 'fixed_n', the neighborhood $N_i$ contains $m - 1$ nearest locations of $s_i$, where $m$ is determined by the parameter neighborhood_size. For more details see Ernst & Haesbroeck, (2017).

After calculating the neighborhoods, the local outliers are generated following Ernst & Haesbroeck, (2017) and Harris et al. (2014) using the steps:

1. Sort the observations from highest to lowest by their principle component analysis (PCA) scores of the first component (PC-1).
2. Set $k$ to be $\alpha N/2$ rounded to nearest integer and select the set of local outlier points $S_{out}$ by finding $k$ observations with the highest PC-1 values and $k$ observations with the lowest PC-1 values under the condition that for all $s_i, s_j \in S_{out}$ it holds that $N_i \neq N_j$.
3. Form sets $X_{large}$, which contains $k$ observations with the largest PC-1 values of outlier points $S_{out}$ and $X_{small}$, which contains $k$ observations with the smallest PC-1 values of outlier points $S_{out}$. Generate the local outliers by swapping $X_{small,i}$ with $X_{large,k+i−1}$, $i = 1, ..., k$. The parameter swap_order defines how the sets $X_{large}$ and $X_{small}$ are ordered.

If the parameter swap_order is 'regular', $X_{small}$ and $X_{large}$ are sorted by PC-1 score from smallest to largest. If the parameter swap_order is 'reverse', $X_{small}$ is sorted from largest to
smallest and \( X_{\text{large}} \) from smallest to largest. If the parameter \( \text{swap\_order} \) is 'random', \( X_{\text{small}} \) and \( X_{\text{large}} \) are in random order.

**Value**

`gen_loc_outl` returns a `data.frame` containing the contaminated fields as \( p \) first columns. The column \( p + 1 \) contains a logical indicator whether the observation is an outlier or not.

**Note**

This function is a modified version of code originally provided by M. Ernst and G. Haesbroeck.

**References**


**See Also**

`gen_glob_outl`

**Examples**

```r
# simulate coordinates
coords <- runif(1000 * 2) * 20
dim(coords) <- c(1000, 2)
coords_df <- as.data.frame(coords)
names(coords_df) <- c("x", "y")

# simulate random field
if (!requireNamespace("gstat", quietly = TRUE)) {
  message("Please install the package gstat to run the example code."
} else {
  library(gstat)
  model_1 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                   model = vgm(psill = 0.025, range = 1, model = "Exp"), nmax = 20)
  model_2 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                   model = vgm(psill = 0.025, range = 1, kappa = 2, model = "Mat"),
                   nmax = 20)
  model_3 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                   model = vgm(psill = 0.025, range = 1, model = "Gau"), nmax = 20)

  field_1 <- predict(model_1, newdata = coords_df, nsim = 1)$sim
  field_2 <- predict(model_2, newdata = coords_df, nsim = 1)$sim
  field_3 <- predict(model_3, newdata = coords_df, nsim = 1)$sim
  field <- cbind(field_1, field_2, field_3)

  # Generate 5 % local outliers to data using radius neighborhoods
```
local_covariance_matrix

Computation of Local Covariance Matrices

**Description**

`local_covariance_matrix` computes local covariance matrices for a random field based on a given set of spatial kernel matrices.

**Usage**

```r
local_covariance_matrix(x, kernel_list, lcov = c('lcov', 'ldiff', 'lcov_norm'), center = TRUE)
```

**Arguments**

- `x` a numeric matrix of dimension \( c(n, p) \) where the \( p \) columns correspond to the entries of the random field and the \( n \) rows are the observations.
- `kernel_list` a list with spatial kernel matrices of dimension \( c(n, n) \). This list is usually computed with the function `spatial_kernel_matrix`.
- `lcov` a string indicating which type of local covariance matrix to use. Either 'lcov' (default) or 'ldiff'.
- `center` logical. If TRUE the data \( x \) is centered prior computing the local covariance matrices. Default is TRUE.

**Details**

Two versions of local covariance matrices are implemented, the argument `lcov` determines which version is used:

- 'lcov':
  \[
  LCov(f) = 1/n \sum_{i,j} f(d_{i,j}) (x(s_i) - \bar{x})(x(s_j) - \bar{x})',
  \]

- 'ldiff':
  \[
  LDiff(f) = 1/n \sum_{i,j} f(d_{i,j}) (x(s_i) - x(s_j))(x(s_i) - x(s_j))',
  \]
local_covariance_matrix

- 'lcov_norm'

\[ LCov^*(f) = \frac{1}{nF_{f,n}^{1/2}} \sum_{i,j} f(d_{i,j})(x(s_i) - \bar{x})(x(s_j) - \bar{x})', \]

with

\[ F_{f,n} = \frac{1}{n} \sum_{i,j} f^2(d_{i,j}). \]

Where \( d_{i,j} \geq 0 \) correspond to the pairwise distances between coordinates, \( x(s_i) \) are the \( p \) random field values at location \( s_i \), \( \bar{x} \) is the sample mean vector, and the kernel function \( f(d) \) determines the locality. The choice 'lcov_norm' is useful when testing for the actual signal dimension of the latent field, see \texttt{sbss_asym} and \texttt{sbss_boot}. The function \texttt{local_covariance_matrix} computes local covariance matrices for a given random field and given spatial kernel matrices, the type of computed local covariance matrices is determined by the argument 'lcov'. If the argument center equals FALSE then the centering in the above formula for \( LCov(f) \) is not carried out. See also \texttt{spatial_kernel_matrix} for details.

Value

\texttt{local_covariance_matrix} returns a list of equal length as the argument \texttt{kernel_list}. Each list entry is a numeric matrix of dimension \( c(p, p) \) corresponding to a local covariance matrix. The list has the attribute 'lcov' which equals the function argument \texttt{lcov}.

References


See Also

\texttt{spatial_kernel_matrix}, \texttt{sbss}

Examples

```r
# simulate coordinates
coords <- runif(1000 * 2) * 20
dim(coords) <- c(1000, 2)
coords_df <- as.data.frame(coords)
names(coords_df) <- c("x", "y")
# simulate random field
if (!requireNamespace("gstat", quietly = TRUE)) {
  message("Please install the package gstat to run the example code."
} else {
  library(gstat)
model_1 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                  model = vgm(psill = 0.025, range = 1, model = "Exp"), nmax = 20)
model_2 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
```
model = vgm(psill = 0.025, range = 1, kappa = 2, model = 'Mat'), nmax = 20)
model_3 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
model = vgm(psill = 0.025, range = 1, model = 'Gau'), nmax = 20)
field_1 <- predict(model_1, newdata = coords_df, nsim = 1)$sim1
field_2 <- predict(model_2, newdata = coords_df, nsim = 1)$sim1
field_3 <- predict(model_3, newdata = coords_df, nsim = 1)$sim1
field <- as.matrix(cbind(field_1, field_2, field_3))

# computing two ring kernel matrices and corresponding local covariance matrices
kernel_params_ring <- c(0, 0.5, 0.5, 2)
ring_kernel_list <-
spatial_kernel_matrix(coords, 'ring', kernel_params_ring)
loc_cov_ring <-
local_covariance_matrix(x = field, kernel_list = ring_kernel_list)

# computing two ring kernel matrices and corresponding local difference matrices
kernel_params_ring <- c(0, 0.5, 0.5, 2)
ring_kernel_list <-
spatial_kernel_matrix(coords, 'ring', kernel_params_ring)
loc_cov_ring <-
local_covariance_matrix(x = field, kernel_list = ring_kernel_list, lcov = 'ldiff')

# computing three ball kernel matrices and corresponding local covariance matrices
kernel_params_ball <- c(0.5, 1, 2)
bball_kernel_list <-
spatial_kernel_matrix(coords, 'ball', kernel_params_ball)
loc_cov_ball <-
local_covariance_matrix(x = field, kernel_list = ball_kernel_list)

# computing three gauss kernel matrices and corresponding local covariance matrices
kernel_params_gauss <- c(0.5, 1, 2)
 gauss_kernel_list <-
spatial_kernel_matrix(coords, 'gauss', kernel_params_gauss)
loc_cov_gauss <-
local_covariance_matrix(x = field, kernel_list = gauss_kernel_list)

---

**local_gss_covariance_matrix**

*Computation of Robust Local Covariance Matrices*

**Description**

`local_gss_covariance_matrix` computes generalized local sign covariance matrices for a random field based on a given set of spatial kernel matrices.

**Usage**

```r
local_gss_covariance_matrix(x, kernel_list, lcov = c('norm', 'winsor', 'qwinsor'), center = TRUE)
```
Arguments

- **x**: a numeric matrix of dimension \(c(n, p)\) where the \(p\) columns correspond to the entries of the random field and the \(n\) rows are the observations.

- **kernel_list**: a list with spatial kernel matrices of dimension \(c(n, n)\). This list is usually computed with the function `spatial_kernel_matrix`.

- **lcov**: a string indicating which type of robust local covariance matrix to use. Either ‘norm’ (default), ‘winsor’ or ‘qwinsor’.

- **center**: logical. If TRUE the data \(x\) is robustly centered prior computing the local covariance matrices. Default is TRUE. See also `white_data`.

Details

Generalized local sign matrices are determined by radial functions \(w(l_i)\), where \(l_i = ||x(s_i) - T(x)||\) and \(T(x)\) is Hettmansperger Randles location estimator (Hettmansperger & Randles, 2002), and kernel functions \(f(d_{i,j})\), where \(d_{i,j} = ||s_i - s_j||\). Generalized local sign covariance (gLSCM) matrix is then calculated as

\[
\text{gLSCM}(f, w) = 1/(n F_{f,n}^{1/2}) \sum_{i,j} f(d_{i,j}) w(l_i) w(l_j) (x(s_i) - T(x))' (x(s_j) - T(x))
\]

with

\[
F_{f,n} = 1/n \sum_{i,j} f^2(d_{i,j}).
\]

Three radial functions \(w(l_i)\) (Raymaekers & Rousseeuw, 2019) are implemented, the parameter \(lcov\) defines which is used:

- ‘norm’:
  \[
w(l_i) = 1/l_i
  \]

- ‘winsor’:
  \[
w(l_i) = Q/l_i
  \]

- ‘qwinsor’:
  \[
w(l_i) = Q^2/l_i^2.
  \]

The cutoff \(Q\) is defined as \(Q = l(h)\), where \(l(h)\) is \(h\)th order statistic of \(\{l_1, \ldots, l_n\}\) and \(h = (n + p + 1)/2\). If the argument center equals FALSE then the centering in the above formula for \(gLSCM(f, w)\) is not carried out. See also `spatial_kernel_matrix` for details.

Value

`local_gss_covariance_matrix` returns a list with two entries:

- **cov_sp_list**: List of equal length as the argument kernel_list. Each list entry is a numeric matrix of dimension \(c(p, p)\) corresponding to a robust local covariance matrix. The list has the attribute ‘lcov’ which equals the function argument \(lcov\).

- **weights**: numeric vector of \(\text{length}(n)\) giving the weights for each observation for the robust local covariance estimation.
References


See Also

spatial_kernel_matrix, robsbss

Examples

# simulate coordinates
coops <- runif(1000 * 2) * 20
dim(coops) <- c(1000, 2)
coops_df <- as.data.frame(coops)
names(coops_df) <- c("x", "y")
# simulate random field
if (!requireNamespace("gstat", quietly = TRUE)) {
    message("Please install the package gstat to run the example code.")
} else {
    library(gstat)
    model_1 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                     model = vgm(psill = 0.025, range = 1, model = 'Exp'), nmax = 20)
    model_2 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                     model = vgm(psill = 0.025, range = 1, kappa = 2, model = 'Mat'),
                     nmax = 20)
    model_3 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                     model = vgm(psill = 0.025, range = 1, model = 'Gau'), nmax = 20)
    field_1 <- predict(model_1, newdata = coops_df, nsim = 1)$sim1
    field_2 <- predict(model_2, newdata = coops_df, nsim = 1)$sim1
    field_3 <- predict(model_3, newdata = coops_df, nsim = 1)$sim1
    field <- cbind(field_1, field_2, field_3)

    # computing two ring kernel matrices and corresponding robust local covariance matrices using 'norm' radial function:
    ring_kernel_params <- c(0, 0.5, 0.5, 2)
    ring_kernel_list <- spatial_kernel_matrix(coops, 'ring', kernel_params_ring)
    loc_cov_ring <- local_gss_covariance_matrix(x = field, kernel_list = ring_kernel_list, lcov = 'norm')

    # computing three ball kernel matrices and corresponding robust local covariance matrices using 'winsor' radial function:
    ball_kernel_params <- c(0.5, 1, 2)
    ball_kernel_list <- spatial_kernel_matrix(coops, 'ball', kernel_params_ball)
    loc_cov_ball <-
local_gss_covariance_matrix(x = field, kernel_list = ball_kernel_list,
   lcov = 'winsor')

# computing three gauss kernel matrices and corresponding
# robust local covariance matrices using 'qwinsor' radial function:
kernel_params_gauss <- c(0.5, 1, 2)
gauss_kernel_list <-
   spatial_kernel_matrix(coords, 'gauss', kernel_params_gauss)
loc_cov_gauss <-
   local_gss_covariance_matrix(x = field, kernel_list = gauss_kernel_list,
      lcov = 'qwinsor')
}

---

plot.sbss  

Plot Method for an Object of Class 'sbss'

Description

plot.sbss is an interface to the standard plot method for the class of the estimated source random field.

Usage

## S3 method for class 'sbss'
plot(x, which = 1:ncol(x$s), ...)

Arguments

x           object of class 'sbss'. Usually result of sbss.
which        a numeric vector indicating which components of the latent field should be plotted.
...          further arguments to the plot method of class(x$s), which is either spplot or plot.

Details

This method calls the corresponding plot method of class(x$s). Either spplot for class(x$s) is SpatialPointsDataFrame or plot.sf for class(x$s) is sf. If x$s is a matrix then it is internally cast to SpatialPointsDataFrame and spplot is used for plotting. Arguments to the corresponding plot functions can be given through ... .

See Also

sbss, spplot, plot.sf
Examples

# simulate coordinates
coords <- runif(1000 * 2) * 20
dim(coords) <- c(1000, 2)
coords_df <- as.data.frame(coords)
names(coords_df) <- c("x", "y")

# simulate random field
if (!requireNamespace("gstat", quietly = TRUE)) {
  message("Please install the package gstat to run the example code."
} else {
  library(gstat)
  model_1 <- gstat(formula = z ~ 1, locations = x + y, dummy = TRUE, beta = 0,
                   model = vgm(psill = 0.025, range = 1, model = "Exp"), nmax = 20)
  model_2 <- gstat(formula = z ~ 1, locations = x + y, dummy = TRUE, beta = 0,
                   model = vgm(psill = 0.025, range = 1, kappa = 2, model = "Mat"),
                   nmax = 20)
  model_3 <- gstat(formula = z ~ 1, locations = x + y, dummy = TRUE, beta = 0,
                   model = vgm(psill = 0.025, range = 1, model = "Gau"), nmax = 20)

  field_1 <- predict(model_1, newdata = coords_df, nsim = 1)$sim1
  field_2 <- predict(model_2, newdata = coords_df, nsim = 1)$sim1
  field_3 <- predict(model_3, newdata = coords_df, nsim = 1)$sim1
  field <- as.matrix(cbind(field_1, field_2, field_3))

  # compute ring kernel matrices
  kernel_parameters <- c(0, 1, 1, 2, 2, 3)
  ring_kernel_list <- spatial_kernel_matrix(coords, "ring", kernel_parameters)

  # apply sbss SpatialPointsDataFrame object
  field_sp <- sp::SpatialPointsDataFrame(coords = coords, data = data.frame(field))
  res_sp <- sbss(field_sp, kernel_list = ring_kernel_list)

  # plot with SpatialPointsDataFrame object
  plot(res_sp)

  # plot with SpatialPointsDataFrame object
  # and additional arguments for spplot function
  plot(res_sp, colorkey = TRUE, as.table = TRUE, cex = 1)

  # apply sbss with sf object
  if (!requireNamespace("sf", quietly = TRUE)) {
    message("Please install the package sf to run the example code."
  } else {
    field_sf <- sf::st_as_sf(data.frame(coords = coords, field),
                             coords = c(1,2))
    res_sf <- sbss(x = field_sf, kernel_list = ring_kernel_list)

    # plot with sf object
    plot(res_sf)

    # and additional arguments for plot.sf function
    plot(res_sf, axes = TRUE, key.pos = 4)
predict.sbss

} } }

predict.sbss

Predict Method for an Object of Class 'sbss'

Description

predict.sbss predicts the estimated source random field on a grid with Inverse Distance Weighting (IDW) and plots these predictions.

Usage

## S3 method for class 'sbss'
predict(object, p = 2, n_grid = 50, which = 1:ncol(object$s), ...)

Arguments

object object of class 'sbss'. Usually result of sbss.
p numeric. The positive power parameter for IDW. Default is 2.
n_grid numeric. Each dimension of the spatial domain is divided by this integer to derive a grid for IDW predictions. Default is 50.
which a numeric vector indicating which components of the latent field should be predicted.
... further arguments to the plot method of class(x$s), which is either spplot or plot.

Details

IDW predictions are made on a grid. The side lengths of the rectangular shaped grid cells are derived by the differences of the rounded maximum and minimum values divided by the n_grid argument for each column of object$coords. Hence, the grid contains a total of n_grid ^ 2 points. The power parameter of the IDW predictions is given by p (default: 2).

The predictions are plotted with the corresponding plot method of class(x$s). Either spplot for class(x$s) is SpatialPointsDataFrame or plot.sf for class(x$s) is sf. If x$s is a matrix then it is internally cast to SpatialPointsDataFrame and spplot is used for plotting. Arguments to the corresponding plot functions can be given through ... as it is done by the method plot.sbss.

Value

The return is dependent on the class of the latent field in the 'sbss' object. If class(object$s) is a matrix then a list with the following entries is returned:

vals_pred_idw a matrix of dimension c(n,p) (when which is default or less than p columns according to the selected components with the which argument) with the IDW predictions of the estimated source random field.
coords_pred_idw

a matrix of dimension c(n, 2) with the grid coordinates for the IDW predictions.

If class(object$s) is SpatialPointsDataFrame or sf then the predicted values and their coordinates are returned as an object of the corresponding class.

The return is invisible.

See Also

sbss, plot.sbss, spplot, plot.sf

Examples

# simulate coordinates
coords <- runif(1000 * 2) * 20
dim(coords) <- c(1000, 2)
coords_df <- as.data.frame(coords)
names(coords_df) <- c("x", "y")

# simulate random field
if (!requireNamespace("gstat", quietly = TRUE)) {
  message("Please install the package gstat to run the example code."
} else {
  library(gstat)
  model_1 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                  model = vgm(psill = 0.025, range = 1, model = 'Exp'), nmax = 20)
  model_2 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                  model = vgm(psill = 0.025, range = 1, kappa = 2, model = 'Mat'),
                  nmax = 20)
  model_3 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                  model = vgm(psill = 0.025, range = 1, model = 'Gau'), nmax = 20)
  field_1 <- predict(model_1, newdata = coords_df, nsim = 1)$sim1
  field_2 <- predict(model_2, newdata = coords_df, nsim = 1)$sim1
  field_3 <- predict(model_3, newdata = coords_df, nsim = 1)$sim1
  field <- as.matrix(cbind(field_1, field_2, field_3))

  # apply sbss with three ring kernels
  kernel_borders <- c(0, 1, 1, 2, 2, 4)
  res_sbss <- sbss(field, coords, "ring", kernel_borders)

  # predict latent fields on grid with default settings
  predict(res_sbss)

  # predict latent fields on grid with custom plotting settings
  predict(res_sbss, colorkey = TRUE, as.table = TRUE, cex = 1)

  # predict latent fields on a 60x60 grid
  predict(res_sbss, n_grid = 60, colorkey = TRUE, as.table = TRUE, cex = 1)

  # predict latent fields with a higher IDW power parameter
  predict(res_sbss, p = 10, colorkey = TRUE, as.table = TRUE, cex = 1)

  # predict latent fields and save the predictions

predict_list <- predict(res_sbss, p = 5, colorkey = TRUE, as.table = TRUE, cex = 1)
}

print.sbss

Print Method for an Object of Class 'sbss'

Description

Prints the estimated unmixing matrix and the diagonalized local covariance matrices for an object of class 'sbss'.

Usage

## S3 method for class 'sbss'
print(x, ...)

Arguments

x object of class 'sbss'. Usually result of sbss.
...
additional arguments for the method print.listof.

See Also

sbss

robsbss

Robust Spatial Blind Source Separation

Description

robsbss is a robust variant of sbss. It estimates the unmixing matrix assuming a spatial blind source separation model by jointly diagonalizing the Hettmansperger-Randles scatter matrix and one/many generalized local sign covariance matrices. These local generalized sign covariance matrices are determined by spatial kernel functions and radial functions. Three types of such kernel functions and three types of radial functions are supported.

Usage

robsbss(x, ...)

## Default S3 method:
robsbss(x, coords, kernel_type = c('ring', 'ball', 'gauss'),
         kernel_parameters, lcov = c('norm', 'winsor', 'qwinsor'),
         ordered = TRUE, kernel_list = NULL, ...)
## S3 method for class 'SpatialPointsDataFrame'
robsbss(x, ...)
## S3 method for class 'sf'
robsbss(x, ...)
Arguments

x either a numeric matrix of dimension \(c(n, p)\) where the \(p\) columns correspond to the entries of the random field and the \(n\) rows are the observations, an object of class `SpatialPointsDataFrame` or an object of class `sf`.

coords a numeric matrix of dimension \(c(n, 2)\) where each row represents the coordinates of a point in the spatial domain. Only needed if \(x\) is a matrix and the argument `kernel_list` is NULL.

kernel_type a string indicating which kernel function to use. Either 'ring' (default), 'ball' or 'gauss'.

kernel_parameters a numeric vector that gives the parameters for the kernel function. At least length of one for 'ball' and 'gauss' or two for 'ring' kernel, see details.

lcov a string indicating which radial function or type of robust local covariance matrix to use. Either 'norm' (default), 'winsor' or 'qwinsor'. See also `local_gss_covariance_matrix` for details.

ordered logical. If TRUE the entries of the latent field are ordered by the sum of squared (pseudo-)eigenvalues of the diagonalized local covariance matrix/matrices. Default is TRUE.

kernel_list a list of spatial kernel matrices with dimension \(c(n,n)\), see details. Usually computed by the function `spatial_kernel_matrix`.

... further arguments for the fast real joint diagonalization algorithm that jointly diagonalizes the local covariance matrices. See details and `frjd`.

Details

`robsbss` is a robust variant of `sbss` which uses Hettmansperger-Randles (HR) location and scatter estimates (Hettmansperger & Randles, 2002) for whitening (see `white_data` for details) and jointly diagonalizes HR scatter matrix and generalized local sign matrices to estimate the unmixing matrix. The generalized local sign matrices are determined by radial functions \(w(l_i)\), where \(l_i = ||x(s_i) - T(x)||\) and \(T(x)\) is HR location estimator, and kernel functions \(f(d_{i,j})\), where \(d_{i,j} = ||s_i - s_j||\). Generalized local sign covariance (gLSCM) matrix is then calculated as

\[
gLSCM(f, w) = 1/(nF_{f,n}^{1/2}) \sum_{i,j} f(d_{i,j})w(l_i)w(l_j)(x(s_i) - T(x))(x(s_j) - T(x))' 
\]

with

\[
F_{f,n} = 1/n \sum_{i,j} f^2(d_{i,j}).
\]

Three radial functions (Raymaekers & Rousseeuw, 2019) \(w(l_i)\) are implemented, the parameter `lcov` defines which is used:

- 'norm':
  \[
  w(l_i) = 1/l_i 
  \]

- 'winsor':
  \[
  w(l_i) = Q/l_i 
  \]
• 'qWinsor':

\[ w(l_i) = \frac{Q^2}{l_i^2}. \]

The cutoff \( Q \) is defined as \( Q = l_{(h)} \), where \( l_{(h)} \) is \( h \)th order statistic of \( \{l_1, \ldots, l_n\} \) and \( h = \frac{n + p + 1}{2} \). In addition, three kernel functions \( f(d) \) are implemented, the parameter kernel_type defines which is used:

1. 'ring': parameters are inner radius \( r_{in} \) and outer radius \( r_{out} \), with \( r_{in} < r_{out} \), and \( r_{in}, r_{out} \geq 0 \):

\[ f(d; r_{in}, r_{out}) = I(r_{in} < d \leq r_{out}) \]

2. 'ball': parameter is the radius \( r \), with \( r \geq 0 \):

\[ f(d; r) = I(d \leq r) \]

3. 'gauss': Gaussian function where 95\% of the mass is inside the parameter \( r \), with \( r \geq 0 \):

\[ f(d; r) = \exp\left(-0.5\Phi^{-1}(0.95)d/r\right)^2. \]

The argument kernel_type determines the used kernel function as presented above, the argument kernel_parameters gives the corresponding parameters for the kernel function. Specifically, if kernel_type equals 'ball' or 'gauss' then kernel_parameters is a numeric vector where each entry corresponds to one parameter. Hence, length(kernel_parameters) local covariance matrices are used. Whereas, if kernel_type equals 'ring', then kernel_parameters must be a numeric vector of even length where subsequently the inner and outer radii must be given (informally: \( c(r_{in1}, r_{out1}, r_{in2}, r_{out2}, \ldots) \)). In that case length(kernel_parameters) / 2 local covariance matrices are used.

robsbss calls spatial_kernel_matrix internally to compute a list of \( c(n, n) \) kernel matrices based on the parameters given, where each entry of those matrices corresponds to \( f(d_{i,j}) \). Alternatively, such a list of kernel matrices can be given directly to the function robsbss via the kernel_list argument. This is useful when robsbss is called numerous times with the same coordinates/kernel functions as the computation of the kernel matrices is then done only once prior the actual robsbss calls. For details see also spatial_kernel_matrix.

If more than one generalized local sign covariance matrix is used robsbss jointly diagonalizes these matrices with the function frjd. ... provides arguments for frjd, useful arguments might be:

- eps: tolerance for convergence.
- maxiter: maximum number of iterations.

Value

robsbss returns a list of class 'sbss' with the following entries:

- s: object of class(x) containing the estimated source random field.
- coords: coordinates of the observations. Is NULL if x was a matrix and the argument kernel_list was not NULL at the robsbss call.
- w: estimated unmixing matrix.
- weights: numeric vector of length(n) giving the weights for each observation for the robust local covariance estimation.
w_inv  inverse of the estimated unmixing matrix.
pevals  (pseudo-)eigenvalues for each latent field entry.
d  matrix of stacked (jointly) diagonalized local covariance matrices with dimension $c(\text{length}(\text{kernel\_parameters}) \times p, p)$ for 'ball' and 'gauss' kernel or $c(\text{(length}(\text{kernel\_parameters}) \div 2) \times p, p)$ for 'ring' kernel.
diags  matrix of dimension $c(\text{length}(\text{kernel\_parameters}), p)$ where the rows contain the diagonal of the diagonalized local autocovariance matrices.
x_mu  robustly estimated column means of $x$.
cov_inv_sqrt  square root of the inverse sample covariance matrix of $x$.

References


See Also

`spatial_kernel_matrix`, `local_gss_covariance_matrix`, `sp`, `sf`, `frjd`

Examples

```r
# simulate coordinates
coords <- runif(1000 * 2) * 20
dim(coords) <- c(1000, 2)
coords_df <- as.data.frame(coords)
names(coords_df) <- c("x", "y")

# simulate random field
if (!requireNamespace("gstat", quietly = TRUE)) {
  message("Please install the package gstat to run the example code."")
} else {
  library(gstat)
  model_1 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                   model = vgm(psill = 0.025, range = 1, model = 'Exp'), nmax = 20)
  model_2 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                   model = vgm(psill = 0.025, range = 1, kappa = 2, model = 'Mat'),
                   nmax = 20)
  model_3 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                   model = vgm(psill = 0.025, range = 1, model = 'Gau'), nmax = 20)
  field_1 <- predict(model_1, newdata = coords_df, nsim = 1)$sim1
  field_2 <- predict(model_2, newdata = coords_df, nsim = 1)$sim1
  field_3 <- predict(model_3, newdata = coords_df, nsim = 1)$sim1
  field <- cbind(field_1, field_2, field_3)
  # Generate 5% local outliers to data
  field_cont <- gen_loc_outl(field, coords, radius = 2,
                             swap_order = "regular")[,1:3]
```
X <- as.matrix(field_cont)

# apply sbss with three ring kernels
kernel_parameters <- c(0, 1, 1, 2, 2, 3)
robsbss_result <-
    robsbss(X, coords, kernel_type = 'ring', kernel_parameters = kernel_parameters)

# print object
print(robsbss_result)

# plot latent field
plot(robsbss_result, colorkey = TRUE, as.table = TRUE, cex = 1)

# predict latent fields on grid
predict(robsbss_result, colorkey = TRUE, as.table = TRUE, cex = 1)

# unmixing matrix
w_unmix <- coef(robsbss_result)

---

sbss  
Spatial Blind Source Separation

Description

sbss estimates the unmixing matrix assuming a spatial blind source separation model by simultaneous/jointly diagonalizing the covariance matrix and one/many local covariance matrices. These local covariance matrices are determined by spatial kernel functions. Three types of such kernel functions are supported.

Usage

sbss(x, ...)

Arguments

x  
either a numeric matrix of dimension c(n, p) where the p columns correspond to the entries of the random field and the n rows are the observations, an object of class SpatialPointsDataFrame or an object of class sf.
coords a numeric matrix of dimension $c(n,2)$ where each row represents the coordinates of a point in the spatial domain. Only needed if $x$ is a matrix and the argument kernel_list is NULL.

kernel_type a string indicating which kernel function to use. Either 'ring' (default), 'ball' or 'gauss'.

kernel_parameters a numeric vector that gives the parameters for the kernel function. At least length of one for 'ball' and 'gauss' or two for 'ring' kernel, see details.

lcov a string indicating which type of local covariance matrix to use. Either 'lcov' (default), 'ldiff' or 'lcov_norm'. See sbss_asym for details on the latter option.

ordered logical. If TRUE the entries of the latent field are ordered by the sum of squared (pseudo)-eigenvalues of the diagonalized local covariance matrix/matrices. Default is TRUE.

kernel_list a list of spatial kernel matrices with dimension $c(n,n)$, see details. Usually computed by the function spatial_kernel_matrix.

rob_whitening logical. If TRUE whitening is carried out with respect to the first spatial scatter matrix and not the sample covariance matrix, see details. Default is FALSE.

... further arguments for the fast real joint diagonalization algorithm that jointly diagonalizes the local covariance matrices. See details and frjd.

Details

Three versions of local covariance matrices are implemented, the argument lcov determines which version is used:

- 'lcov':
  \[ LCov(f) = \frac{1}{n} \sum_{i,j} f(d_{i,j})(x(s_i) - \bar{x})(x(s_j) - \bar{x})', \]

- 'ldiff':
  \[ LDiff(f) = \frac{1}{n} \sum_{i,j} f(d_{i,j})(x(s_i) - x(s_j))(x(s_i) - x(s_j))', \]

- 'lcov_norm':
  \[ LCov^*(f) = \frac{1}{nF_{f,n}^{1/2}} \sum_{i,j} f(d_{i,j})(x(s_i) - \bar{x})(x(s_j) - \bar{x})', \]

with

\[ F_{f,n} = \frac{1}{n} \sum_{i,j} f^2(d_{i,j}). \]

Where $d_{i,j} \geq 0$ correspond to the pairwise distances between coordinates, $x(s_i)$ are the $p$ random field values at location $s_i$, $\bar{x}$ is the sample mean vector, and the kernel function $f(d)$ determines the locality. The choice 'lcov_norm' is useful when testing for the actual signal dimension of the latent field, see sbss_asym and sbss_boot. LDiff matrices are supposed to be more robust when the random field shows a smooth trend. The following kernel functions are implemented and chosen with the argument kernel_type:
'ring': parameters are inner radius $r_{in}$ and outer radius $r_{out}$, with $r_{in} < r_{out}$, and $r_{in}, r_{out} \geq 0$:

$$f(d; r_{in}, r_{out}) = I(r_{in} < d \leq r_{out})$$

'ball': parameter is the radius $r$, with $r \geq 0$:

$$f(d; r) = I(d \leq r)$$

'gauss': Gaussian function where 95% of the mass is inside the parameter $r$, with $r \geq 0$:

$$f(d; r) = \exp(-0.5(\Phi^{-1}(0.95)d/r)^2)$$

The argument kernel_type determines the used kernel function as presented above, the argument kernel_parameters gives the corresponding parameters for the kernel function. Specifically, if kernel_type equals 'ball' or 'gauss' then kernel_parameters is a numeric vector where each entry corresponds to one parameter. Hence, length(kernel_parameters) local covariance matrices are used. Whereas, if kernel_type equals 'ring', then kernel_parameters must be a numeric vector of even length where subsequently the inner and outer radii must be given (informally: c(r_in1, r_out1, r_in2, r_out2, ...)). In that case length(kernel_parameters) / 2 local covariance matrices are used.

Internally, sbss calls spatial_kernel_matrix to compute a list of c(n,n) kernel matrices based on the parameters given, where each entry of those matrices corresponds to one parameter. Hence, length(kernel_parameters) local covariance matrices are used. Whereas, if kernel_type equals 'ring', then kernel_parameters must be a numeric vector of even length where subsequently the inner and outer radii must be given (informally: c(r_in1, r_out1, r_in2, r_out2, ...)). In that case length(kernel_parameters) / 2 local covariance matrices are used.

rob_whitening determines which scatter is used for the whitening step. If TRUE, whitening is carried out with respect to the scatter matrix defined by the lcov argument, where the kernel function is given by the argument kernel_type and the parameters correspond to the first occurring in the argument kernel_parameters. Therefore, at least two different kernel parameters need to be given. Note that only LDiff(f) matrices are positive definite, hence whitening with 'lcov' is likely to produce an error. If the argument is FALSE, whitening is carried out with respect to the usual sample covariance matrix. sbss internally calls white_data.

If more than one local covariance matrix is used sbss jointly diagonalizes these matrices with the function frjd. ... provides arguments for frjd, useful arguments might be:

- eps: tolerance for convergence.
- maxiter: maximum number of iterations.

Value

sbss returns a list of class 'sbss' with the following entries:

- s: object of class(x) containing the estimated source random field.
- coords: coordinates of the observations. Is NULL if x was a matrix and the argument kernel_list was not NULL at the sbss call.
- w: estimated unmixing matrix.
- w_inv: inverse of the estimated unmixing matrix.
pevals  (pseudo-)eigenvalues for each latent field entry.
d  matrix of stacked (jointly) diagonalized local covariance matrices with dimension c(length(kernel_parameters)*p,p) for 'ball' and 'gauss' kernel or c( (length(kernel_parameters) / 2)*p,p) for 'ring' kernel.
diags matrix of dimension c(length(kernel_parameters),p) where the rows contain the diagonal of the diagonalized local autocovariance matrices.
x_mu  columnmeans of x.
cov_inv_sqrt square root of the inverse sample covariance matrix of x.

References


See Also

spatial_kernel_matrix, local_covariance_matrix, sp, sf, frjd

Examples

```r
# simulate coordinates
coords <- runif(1000 * 2) * 20
dim(coords) <- c(1000, 2)
coords_df <- as.data.frame(coords)
names(coords_df) <- c("x", "y")

# simulate random field
if (!requireNamespace("gstat", quietly = TRUE)) {
  message("Please install the package gstat to run the example code.")
} else {
  library(gstat)
  model_1 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                    model = vgm(psill = 0.025, range = 1, model = 'Exp'), nmax = 20)
  model_2 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                    model = vgm(psill = 0.025, range = 1, kappa = 2, model = 'Mat'),
                    nmax = 20)
  model_3 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                    model = vgm(psill = 0.025, range = 1, model = 'Gau'), nmax = 20)
  field_1 <- predict(model_1, newdata = coords_df, nsim = 1)$sim1
  field_2 <- predict(model_2, newdata = coords_df, nsim = 1)$sim1
  field_3 <- predict(model_3, newdata = coords_df, nsim = 1)$sim1
  field <- as.matrix(cbind(field_1, field_2, field_3))
```
# apply sbss with three ring kernels
kernel_parameters <- c(0, 1, 1, 2, 2, 3)
sbss_result <-
  sbss(field, coords, kernel_type = 'ring', kernel_parameters = kernel_parameters)

# print object
print(sbss_result)

# plot latent field
plot(sbss_result, colorkey = TRUE, as.table = TRUE, cex = 1)

# predict latent fields on grid
predict(sbss_result, colorkey = TRUE, as.table = TRUE, cex = 1)

# unmixing matrix
w_unmix <- coef(sbss_result)

# apply the same sbss with a kernel list
kernel_list <- spatial_kernel_matrix(coords, kernel_type = 'ring', kernel_parameters)
sbss_result_k <- sbss(field, kernel_list = kernel_list)

# apply sbss with three ring kernels and local difference matrices
sbss_result_ldiff <-
  sbss(field, coords, kernel_type = 'ring',
       kernel_parameters = kernel_parameters, lcov = 'ldiff')

---

sbss_asympt

Asymptotic Test for the White Noise Dimension in a Spatial Blind Source Separation Model

Description

sbss_asympt uses asymptotic theory for the spatial blind source separation (SBSS) methodology to test if the last \( p - q \) entries of the latent random field are white noise assuming that the \( p \)-variate observed random field follows a SBSS model.

Usage

sbss_asympt(x, ...)

## Default S3 method:
sbss_asympt(x, coords, q, kernel_parameters,
            kernel_list = NULL, ...)

## S3 method for class 'SpatialPointsDataFrame'
sbss_asympt(x, ...)

## S3 method for class 'sf'
sbss_asympt(x, ...)
Arguments

- **x**: either a numeric matrix of dimension \(c(n, p)\) where the \(p\) columns correspond to the entries of the random field and the \(n\) rows are the observations, an object of class `SpatialPointsDataFrame` or an object of class `sf`.

- **coords**: a numeric matrix of dimension \(c(n, 2)\) where each row represents the coordinates of a point in the spatial domain. Only needed if \(x\) is a matrix and the argument `kernel_list` is `NULL`.

- **q**: an integer between \(0\) and \(p - 1\) specifying the number of hypothetical signal components (null hypothesis) in the latent random field.

- **kernel_parameters**: a numeric vector that gives the parameters for the ring kernel function. At least length of two, see details.

- **kernel_list**: a list of spatial kernel matrices with dimension \(c(n, n)\), see details. Usually computed by the function `spatial_kernel_matrix`.

- **...**: further arguments for the fast real joint diagonalization algorithm that jointly diagonalizes the local covariance matrices. See details and `frjd`.

Details

This function uses the SBSS methodology in conjunction with local covariance matrices based on ring kernel functions to estimate the \(p\)-variate latent random field \(s = x^{wh}w\), where \(x^{wh}\) is the whitened version of the data and \(w\) is the estimated unmixing matrix. The considered (adapted) local covariance matrices write as

\[
LCov^* = 1/n F_n^{1/2} \sum_{i,j} I(r_i < d_{i,j} \leq r_o) (x(s_i) - \bar{x})(x(s_j) - \bar{x})'
\]

with

\[
F_n = 1/n \sum_{i,j} I(r_i < d_{i,j} \leq r_o).
\]

Where \(d_{i,j} \geq 0\) correspond to the pairwise distances between coordinates, \(x(s_i)\) are the \(p\) random field values at location \(s_i\) (which is the \(i\)-th row of the argument \(x\) and the location corresponds to the \(i\)-th row of the argument `coords`) and \(\bar{x}\) is the sample mean vector. The function argument `kernel_parameters` determines the parameters of the used ring kernel functions or alternatively a list of kernel matrices can be given with the argument `kernel_list`, see `sbss` for details.

The null hypothesis specified with the argument `q` states that the last \(p - q\) components of the estimated latent field are white noise. The method orders the components of the latent field by the order of the decreasing sums of squares of the corresponding (pseudo-)eigenvalues of the local covariance matrices produced by the joint diagonalization algorithm (or the eigendecomposition if only one local covariance matrix is used). Under the null the lower right \((p - q) \times (p - q)\) block matrices of the jointly diagonalized local covariance matrices equal zero matrices. Therefore, the sum of their squared norms \(m\) is used as test statistic.

This function conducts the hypothesis test using the asymptotic null distribution of \(m\), a chi-squared distribution with \(k(p - q)(p - q + 1)/2\) degrees of freedom \((k\) is the number jointly diagonalized local covariance matrices).

If more than one local covariance matrix is used `sbss_asym` jointly diagonalizes these matrices with the function `frjd`. ... provides arguments for `frjd`, useful arguments might be:


- eps: tolerance for convergence.
- maxiter: maximum number of iterations.

**Value**

`sbss_asym` returns a list of class 'sbss_test' inheriting from the classes 'htest' and 'sbss' with the following entries:

- alternative: a string containing the alternative hypothesis.
- method: a string which indicates which test methods was used.
- data.name: a string specifying the name of the used data.
- statistic: the value of the test statistic.
- parameters: degrees of freedom for the asymptotic chi-squared distribution of the test statistic under the null hypothesis.
- p.value: the p-value of the test.
- s: object of class(x) containing the estimated source random field.
- coords: coordinates of the observations. Is NULL if x was a matrix and the argument kernel_list was not NULL at the `sbss_asym` call.
- w: estimated unmixing matrix.
- w_inv: inverse of the estimated unmixing matrix.
- d: matrix of stacked (jointly) diagonalized local covariance matrices with dimension c((length(kernel_parameters) / 2)*p,p).
- x_mu: columnmeans of x.
- cov_inv_sqrt: square root of the inverse sample covariance matrix of x.

**References**


**See Also**

`sbss`, `spatial_kernel_matrix`, `local_covariance_matrix`, `sp`, `sf`, `frjd`

**Examples**

```r
# simulate coordinates
n <- 1000
coords <- runif(n * 2) * 20
dim(coords) <- c(n, 2)
coords_df <- as.data.frame(coords)
names(coords_df) <- c("x", "y")
# simulate random field
if (!requireNamespace('gstat', quietly = TRUE)) {
  message('Please install the package gstat to run the example code."
} else {
  library(gstat)
```
model_1 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
model = vgm(psill = 0.025, range = 1, model = 'Exp'), nmax = 20)
model_2 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
model = vgm(psill = 0.025, range = 1, kappa = 2, model = 'Mat'),
nmax = 20)
field_1 <- predict(model_1, newdata = coords_df, nsim = 1)$sim1
field_2 <- predict(model_2, newdata = coords_df, nsim = 1)$sim1
field_3 <- rnorm(n)
field_4 <- rnorm(n)
latent_field <- cbind(as.matrix(cbind(field_1, field_2)), field_3, field_4)
mixing_matrix <- matrix(rnorm(16), 4, 4)
observed_field <- latent_field %*% t(mixing_matrix)

# apply the asymptotic test for a hypothetical latent white noise dimension of q
# q can lie between 0 and 3 in this case
# using one ring kernel function and the null hypothesis q = 1
asymp_res_1 <- sbss_asymp(observed_field, coords, q = 1, kernel_parameters = c(0, 1))

# using two ring kernel functions and the null hypothesis q = 3
asymp_res_2 <- sbss_asymp(observed_field, coords, q = 3, kernel_parameters = c(0, 1, 1, 2))

# the result is of class sbss_test which is inherited from htest and sbss
# print object (print method for an object of class htest)
print(asymp_res_1)
print(asymp_res_2)

# plot latent field (plot method for an object of class sbss)
plot(asymp_res_1, colorkey = TRUE, as.table = TRUE, cex = 1)

# predict latent fields on grid (predict method for an object of class sbss)
predict(asymp_res_1, colorkey = TRUE, as.table = TRUE, cex = 1)

# unmixing matrix (coef method for an object of class sbss)
w_unmix <- coef(asymp_res_1)
}

---

**Description**

sbss_boot uses bootstrap tests for the spatial blind source separation (SBSS) methodology to test if the last \( p - q \) entries of the latent random field are white noise assuming that the \( p \)-variate observed random field follows a SBSS model.
Usage

```r
sbss_boot(x, ...)

## Default S3 method:
sbss_boot(x, coords, q, kernel_parameters,
        boot_method = c("permute", "parametric"),
        n_boot = 200, kernel_list = NULL, ...)
## S3 method for class 'SpatialPointsDataFrame'
sbss_boot(x, ...)
## S3 method for class 'sf'
sbss_boot(x, ...)
```

Arguments

- `x` either a numeric matrix of dimension `c(n, p)` where the `p` columns correspond to the entries of the random field and the `n` rows are the observations, an object of class `SpatialPointsDataFrame` or an object of class `sf`.
- `coords` a numeric matrix of dimension `c(n,2)` where each row represents the coordinates of a point in the spatial domain. Only needed if `x` is a matrix and the argument `kernel_list` is `NULL`.
- `q` an integer between 0 and `p - 1` specifying the number of hypothetical signal components (null hypothesis) in the latent random field.
- `kernel_parameters` a numeric vector that gives the parameters for the ring kernel function. At least length of two, see details.
- `boot_method` a string indicating which bootstrap strategy is used, see details. Either 'permute' (default) or 'parametric'.
- `n_boot` positive integer specifying the number of bootstrap samples. Default is 200.
- `kernel_list` a list of spatial kernel matrices with dimension `c(n,n)`, see details. Usually computed by the function `spatial_kernel_matrix`.
- `...` further arguments for the fast real joint diagonalization algorithm that jointly diagonalizes the local covariance matrices. See details and `frjd`.

Details

This function uses the SBSS methodology in conjunction with local covariance matrices based on ring kernel functions to estimate the `p`-variate latent random field `s = x^{wh} u`, where `x^{wh}` is the whitened version of the data and `u` is the estimated unmixing matrix. The considered (adapted) local covariance matrices write as

\[
LCov^* = 1/(n F_n^{1/2}) \sum_{i,j} I(r_i < d_{i,j} \leq r_o) \langle x(s_i) - \bar{x} \rangle \langle x(s_j) - \bar{x} \rangle'
\]

with

\[
F_n = 1/n \sum_{i,j} I(r_i < d_{i,j} \leq r_o).
\]
Where $d_{i,j} \geq 0$ correspond to the pairwise distances between coordinates, $x(s_i)$ are the $p$ random field values at location $s_i$ (which is the $i$-th row of the argument $x$ and the location corresponds to the $i$-th row of the argument $coords$) and $\bar{x}$ is the sample mean vector. The function argument `kernel_parameters` determines the parameters of the used ring kernel functions or alternatively a list of kernel matrices can be given with the argument `kernel_list`, see `sbss` for details.

The null hypothesis specified with the argument $q$ states that the last $p-q$ components of the estimated latent field are white noise. The method orders the components of the latent field by the order of the decreasing sums of squares of the corresponding (pseudo-)eigenvalues of the local covariance matrices produced by the joint diagonalization algorithm (or the eigendecomposition if only one local covariance matrix is used). Under the null the lower right $(p-q) \times (p-q)$ block matrices of the jointly diagonalized local covariance matrices equal zero matrices. Therefore, the sum of their squared norms $m$ is used as test statistic for the bootstrap based inference methods described below.

1. Compute the test statistic $m$ based on the original data $x$.
2. The estimated latent field $s$ (its dimension is $c(n,p)$) is split into the signal part (first $q$ columns) and the white noise part (last $p-q$ columns).
3. Replace the noise part by a bootstrap sample drawn based on one of the two strategies described below.
4. Recombine the signal part and resampled noise part by concatenating the columns leading to $s^{bs}$ and back-transform it by $x^{bs} = s^{bs} w^{-1}$.
5. Compute the test statistic $m^{bs}$ based on $x^{bs}$.
6. Repeat Step 2 - 5 for a total amount of $n_{\text{boot}}$ times (default is 200) and the p-value of the bootstrap test is computed by

$$\frac{\text{sum}(m > m^{bs}) + 1}{n_{\text{boot}} + 1}.$$ 

The argument `boot_method` (default is "permute") specifies the used resample strategy. The two following strategies are implemented:

- `boot_method = "permute"`: This strategy is non-parametric. It draws each bootstrap sample from the vector of all $n(p-q)$ observed hypothetical white noise observations.
- `boot_method = "parametric"`: This is parametric. Each bootstrap sample is drawn independently and identically from the standard normal distribution.

If more than one local covariance matrix is used `sbss_boot` jointly diagonalizes these matrices with the function `frjd`. ... provides arguments for `frjd`, useful arguments might be:

- `eps`: tolerance for convergence.
- `maxiter`: maximum number of iterations.

**Value**

`sbss_boot` returns a list of class `sbss_test` inheriting from the classes `htest` and `sbss` with the following entries:

- `alternative`: a string containing the alternative hypothesis.
- `method`: a string which indicates which test methods was used.
sbss_boot

data.name a string specifying the name of the used data.
statistic the value of the test statistic.
parameters a integer specifying the number of generated bootstrap samples (the value of the argument n_boot).
p.value the p-value of the test.
s object of class(x) containing the estimated source random field.
coords coordinates of the observations. Is NULL if x was a matrix and the argument kernel_list was not NULL at the sbss_boot call.
w estimated unmixing matrix.
w_inv inverse of the estimated unmixing matrix.
d matrix of stacked (jointly) diagonalized local covariance matrices with dimension c((length(kernel_parameters) / 2)*p,p).
x_mu columnmeans of x.
cov_inv_sqrt square root of the inverse sample covariance matrix of x.

References


See Also

sbss, spatial_kernel_matrix, local_covariance_matrix, sp, sf, frjd

Examples

# simulate coordinates
n <- 1000
coords <- runif(n * 2) * 20
dim(coords) <- c(n, 2)
coords_df <- as.data.frame(coords)
names(coords_df) <- c("x", "y")
# simulate random field
if (!requireNamespace("gstat", quietly = TRUE)) {
  message("Please install the package gstat to run the example code.")
} else {
  library(gstat)
  model_1 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
  model = vgm(psill = 0.025, range = 1, model = 'Exp'), nmax = 20)
  model_2 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
  model = vgm(psill = 0.025, range = 1, kappa = 2, model = 'Mat'),
  nmax = 20)
  field_1 <- predict(model_1, newdata = coords_df, nsim = 1)$sim1
  field_2 <- predict(model_2, newdata = coords_df, nsim = 1)$sim1
  field_3 <- rnorm(n)
  field_4 <- rnorm(n)
  latent_field <- cbind(as.matrix(cbind(field_1, field_2)), field_3, field_4)
mixing_matrix <- matrix(rnorm(16), 4, 4)
observed_field <- latent_field %*% t(mixing_matrix)

# apply the bootstrap tests for a hypothetical latent white noise dimension of q
# q can lie between 0 and 3 in this case
# using one ring kernel function with the permute strategy
# and the null hypothesis q = 1
boot_res_1 <-
sbss.boot(observed_field, coords, q = 1, kernel_parameters = c(0, 1),
          boot_method = 'permute', n_boot = 100)

# using two one ring kernel function with the parametric strategy
# and the null hypothesis q = 3
boot_res_2 <-
sbss.boot(observed_field, coords, q = 3, kernel_parameters = c(0, 1, 1, 2),
          boot_method = 'parametric', n_boot = 100)

# the result is of class sbss_test which is inherited from htest and sbss
# print object (print method for an object of class htest)
print(boot_res_1)
print(boot_res_2)

# plot latent field (plot method for an object of class sbss)
plot(boot_res_1, colorkey = TRUE, as.table = TRUE, cex = 1)

# predict latent fields on grid (predict method for an object of class sbss)
predict(boot_res_1, colorkey = TRUE, as.table = TRUE, cex = 1)

# unmixing matrix (coef method for an object of class sbss)
w_unmix <- coef(boot_res_1)

---

**snss_jd**

**Spatial Non-Stationary Source Separation Joint Diagonalization**

**Description**

snss_jd estimates the unmixing matrix assuming a spatial non-stationary source separation model implying non-constant covariance by jointly diagonalizing at least two covariance matrices computed for corresponding different sub-domains.

**Usage**

```r
snss_jd(x, ...)
```

## Default S3 method:

```r
snss_jd(x, coords, n_block, ordered = TRUE, ...)
```

## S3 method for class 'list'

```r
snss_jd(x, coords, ordered = TRUE, ...)
```
## S3 method for class 'SpatialPointsDataFrame'
snss_jd(x, ...)

## S3 method for class 'sf'
snss_jd(x, ...)

### Arguments

- **x**: either a numeric matrix of dimension \(c(n, p)\) where the \(p\) columns correspond to the entries of the random field and the \(n\) rows are the observations, a list of length \(K\) defining the subdivision of the domain, an object of class \(sf\) or an object of class \(SpatialPointsDataFrame\).

- **coords**: a numeric matrix of dimension \(c(n, 2)\) when \(x\) is a matrix where each row represents the sample location of a point in the spatial domain or a list of length \(K\) if \(x\) is a list which defines the subdivision of the domain. Not needed otherwise.

- **n_block**: an integer defining the subdivision of the domain. See details.

- **ordered**: logical. If \(TRUE\) the entries of the latent field are ordered by the sum of squared pseudo-eigenvalues of the diagonalized sub-domain covariance matrices. Default is \(TRUE\).

- **...**: further arguments for the fast real joint diagonalization algorithm that jointly diagonalizes the sub-domain covariance matrices. See details and \(frjd\).

### Details

This function assumes that the random field \(x\) is formed by

\[ x(t) = As(t) + b, \]

where \(A\) is the deterministic \(p \times p\) mixing matrix, \(b\) is the \(p\)-dimensional location vector, \(x\) is the observable \(p\)-variate random field given by the argument \(x\), \(t\) are the spatial locations given by the argument \(coords\) and \(s\) is the latent \(p\)-variate random field assumed to consist of uncorrelated entries that have zero mean but non-constant variances. This function aims to recover \(s\) by

\[ W(x(t) - \bar{x}), \]

where \(W\) is the \(p \times p\) unmixing matrix and \(\bar{x}\) is the sample mean. The function does this by splitting the given spatial domain into \(n_{\text{block}}^2\) equally sized rectangular sub-domains and jointly diagonalizing the corresponding covariance matrices for all sub-domains.

Alternatively the domain subdivision can be defined by providing lists of length \(K\) for the arguments \(x\) and \(coords\) where the first list entries correspond to the values and coordinates of the first sub-domain and the second entries to the values and coordinates of the second sub-domain, etc..

\(snss_jd\) jointly diagonalizes the covariance matrices for each sub-domain with the function \(frjd\). ... provides arguments for \(frjd\), useful arguments might be:

- **eps**: tolerance for convergence.
- **maxiter**: maximum number of iterations.
Value

Similarly as `sbss` the function `snss_jd` returns a list of class `snss` and `sbss` with the following entries:

- **s**: object of class(x) containing the estimated source random field.
- **coords**: coordinates of the observations. Only given if x is a matrix or list.
- **w**: estimated unmixing matrix.
- **w_inv**: inverse of the estimated unmixing matrix.
- **d**: matrix of stacked (jointly) diagonalized sub-domain covariance matrices with dimension \( c(n_{\text{block}}^2 \times p, p) \) or \( c(K \times p, p) \) if x and coords are lists of length K.
- **x_mu**: columnmeans of x.
- **cov_inv_sqrt**: square root of the inverse sample covariance matrix of x.

References


See Also

`sbss`, `sp`, `sf`

Examples

```r
# simulate coordinates
n <- 1000
coords <- runif(n * 2) * 20
dim(coords) <- c(n, 2)

# simulate random field
field_1 <- rnorm(n)
field_2 <- 2 * sin(pi / 20 * coords[, 1]) * rnorm(n)
field_3 <- rnorm(n) * (coords[, 1] < 10) + rnorm(n, 0, 3) * (coords[, 1] >= 10)
latent_field <- cbind(field_1, field_2, field_3)
mixing_matrix <- matrix(rnorm(9), 3, 3)
observed_field <- latent_field

observed_field_sp <- sp::SpatialPointsDataFrame(coords = coords,
                                               data = data.frame(observed_field))
sp::spplot(observed_field_sp, colorkey = TRUE, as.table = TRUE, cex = 1)

# apply snss_jd with 4 sub-domains
res_4 <- snss_jd(observed_field, coords, n_block = 2)
JADE:::MD(W.hat = coef(res_4), A = mixing_matrix)

# apply snss_jd with 9 sub-domains
res_9 <- snss_jd(observed_field, coords, n_block = 3)
```
JADE::MD(W.hat = coef(res_9), A = mixing_matrix)
cor(res_9$s, latent_field)

# print object
print(res_4)

# plot latent field
plot(res_4, colorkey = TRUE, as.table = TRUE, cex = 1)

# predict latent fields on grid
predict(res_4, colorkey = TRUE, as.table = TRUE, cex = 1)

# unmixing matrix
w_unmix <- coef(res_4)

# apply snss_jd with SpatialPointsDataFrame object
res_4_sp <- snss_jd(observed_field_sp, n_block = 2)

# apply with list arguments
# first axis split by 5
# second axis split by 10
# results in 4 sub-domains
flag_x <- coords[, 1] < 5
flag_y <- coords[, 2] < 10
coords_list <- list(coords[flag_x & flag_y, ],
                     coords[!flag_x & flag_y, ],
                     coords[flag_x & !flag_y, ],
                     coords[!flag_x & !flag_y, ])
field_list <- list(observed_field[flag_x & flag_y, ],
                   observed_field[!flag_x & flag_y, ],
                   observed_field[flag_x & !flag_y, ],
                   observed_field[!flag_x & !flag_y, ])
plot(coords, col = 1)
points(coords_list[[2]], col = 2)
points(coords_list[[3]], col = 3)
points(coords_list[[4]], col = 4)

res_list <- snss_jd(x = field_list,
                    coords = coords_list)
plot(res_list, colorkey = TRUE, as.table = TRUE, cex = 1)
JADE::MD(W.hat = coef(res_list), A = mixing_matrix)

---

**snss_sd**

**Spatial Non-Stationary Source Separation Simultaneous Diagonalization**

**Description**

`snss_sd` estimates the unmixing matrix assuming a spatial non-stationary source separation model.
implying non-constant covariance by simultaneously diagonalizing two covariance matrices com-
punted for two corresponding different sub-domains.

Usage

```r
snss_sd(x, ...)
```

### Default S3 method:

```r
snss_sd(x, coords, direction = c('x', 'y'),
        ordered = TRUE, ...)
```

### S3 method for class 'list'

```r
snss_sd(x, coords, ordered = TRUE, ...)
```

### S3 method for class 'SpatialPointsDataFrame'

```r
snss_sd(x, ...)
```

### S3 method for class 'sf'

```r
snss_sd(x, ...)
```

Arguments

- **x**: either a numeric matrix of dimension \(c(n, p)\) where the \(p\) columns correspond to the entries of the random field and the \(n\) rows are the observations, a list of length two defining the subdivision of the domain, an object of class `sf` or an object of class `SpatialPointsDataFrame`.

- **coords**: a numeric matrix of dimension \(c(n, 2)\) when \(x\) is a matrix where each row represents the sample location of a point in the spatial domain or a list of length two if \(x\) is a list which defines the subdivision of the domain. Not needed otherwise.

- **direction**: a string indicating on which coordinate axis the domain is halved. Either `’x’` (default) or `’y’`.

- **ordered**: logical. If `TRUE` the entries of the latent field are ordered according to the de-
creasingly ordered eigenvalues. Default is `TRUE`.

- **...**: further arguments to be passed to or from methods.

Details

This function assumes that the random field \(x\) is formed by

\[
x(t) = As(t) + b,
\]

where \(A\) is the deterministic \(p \times p\) mixing matrix, \(b\) is the \(p\)-dimensional location vector, \(x\) is the observable \(p\)-variate random field given by the argument \(x\), \(t\) are the spatial locations given by the argument `coords` and \(s\) is the latent \(p\)-variate random field assumed to consist of uncorrelated entries that have zero mean but non-constant variances. This function aims to recover \(s\) by

\[
W(x(t) - \bar{x}),
\]

where \(W\) is the \(p \times p\) unmixing matrix and \(\bar{x}\) is the sample mean. The function does this by splitting the given spatial domain in half according to the first coordinate (argument `direction` equals `’x’`) or the second coordinate (argument `direction` equals `’y’`) and simultaneously diagonalizing the sample covariance matrices for each of the two sub-domains.
Alternatively the domain subdivision can be defined by providing lists of length two for the arguments `x` and `coords` where the first list entries correspond to the values and coordinates of the first sub-domain and the second entries to the values and coordinates of the second sub-domain.

**Value**

Similarly as `sbss` the function `snss_sd` returns a list of class `’snss’` and `’sbss’` with the following entries:

- `s` object of class `class(x)` containing the estimated source random field.
- `coords` coordinates of the observations. Only given if `x` is a matrix or list.
- `w` estimated unmixing matrix.
- `w_inv` inverse of the estimated unmixing matrix.
- `d` diagonal matrix containing the eigenvalues of the eigendecomposition.
- `x_mu` columnmeans of `x`.
- `cov_inv_sqrt` square root of the inverse sample covariance matrix for the first sub-domain.

**References**


**See Also**

`sbss`, `sp`, `sf`

**Examples**

```r
# simulate coordinates
n <- 1000
coords <- runif(n * 2) * 20
dim(coords) <- c(n, 2)

# simulate random field
field_1 <- rnorm(n)
field_2 <- 2 * sin(pi / 20 * coords[, 1]) * rnorm(n)
field_3 <- rnorm(n) * (coords[, 1] < 10) + rnorm(n, 0, 3) * (coords[, 1] >= 10)
latent_field <- cbind(field_1, field_2, field_3)
mixing_matrix <- matrix(rnorm(9), 3, 3)
observed_field <- latent_field %*% t(mixing_matrix)
observed_field_sp <- sp::SpatialPointsDataFrame(coords = coords,
                                           data = data.frame(observed_field))
sp::spplot(observed_field_sp, colorkey = TRUE, as.table = TRUE, cex = 1)

# apply snss_sd with split in x
res_x <- snss_sd(observed_field, coords, direction = 'x')
JADE::MD(W.hat = coef(res_x), A = mixing_matrix)
```
# apply snss_sd with split in y
# should be much worse as field shows only variation in x
res_y <- snss_sd(observed_field, coords, direction = 'y')
JADE::MD(W.hat = coef(res_y), A = mixing_matrix)

# print object
print(res_x)

# plot latent field
plot(res_x, colorkey = TRUE, as.table = TRUE, cex = 1)

# predict latent fields on grid
predict(res_x, colorkey = TRUE, as.table = TRUE, cex = 1)

# unmixing matrix
w_unmix <- coef(res_x)

# apply snss_sd with SpatialPointsDataFrame object
res_x_sp <- snss_sd(observed_field_sp, direction = 'x')

# apply with list arguments
# first axis split by 5
flag_coords <- coords[, 1] < 5
coords_list <- list(coords[flag_coords, ],
                     coords[!flag_coords, ])
field_list <- list(observed_field[flag_coords, ],
                   observed_field[!flag_coords, ])
plot(coords, col = flag_coords + 1)
res_list <- snss_sd(x = field_list,
                    coords = coords_list)
plot(res_list, colorkey = TRUE, as.table = TRUE, cex = 1)
JADE::MD(W.hat = coef(res_list), A = mixing_matrix)

---

**snss_sjd**

*Spatial Non-Stationary Source Separation Spatial Joint Diagonalization*

**Description**

*snss_sjd* estimates the unmixing matrix assuming a spatial non-stationary source separation model implying non-constant (spatial) covariance by jointly diagonalizing several covariance and/or spatial covariance matrices computed for a subdivision of the spatial domain into at least two subdomains.

**Usage**

```r
snss_sjd(x, ...)```

## Default S3 method:
snss_sjd(x, coords, n_block, kernel_type = c('ring', 'ball', 'gauss'),
    kernel_parameters, with_cov = TRUE, lcov = c('lcoov', 'ldiff', 'lcoov_norm'),
    ordered = TRUE, ...)
## S3 method for class 'list'
snss_sjd(x, coords, kernel_type = c('ring', 'ball', 'gauss'),
    kernel_parameters, with_cov = TRUE, lcov = c('lcoov', 'ldiff', 'lcoov_norm'),
    ordered = TRUE, ...)
## S3 method for class 'SpatialPointsDataFrame'
snss_sjd(x, ...)
## S3 method for class 'sf'
snss_sjd(x, ...)

### Arguments

- **x**: either a numeric matrix of dimension c(n, p) where the p columns correspond to the entries of the random field and the n rows are the observations, a list of length K defining the subdivision of the domain, an object of class sf or an object of class SpatialPointsDataFrame.
- **coords**: a numeric matrix of dimension c(n,2) when x is a matrix where each row represents the sample location of a point in the spatial domain or a list of length K if x is a list which defines the subdivision of the domain. Not needed otherwise.
- **n_block**: either be an integer defining the subdivision of the domain, 'x' or 'y'. See details.
- **kernel_type**: a string indicating which kernel function to use. Either 'ring' (default), 'ball' or 'gauss'.
- **kernel_parameters**: a numeric vector that gives the parameters for the kernel function. At least length of one for 'ball' and 'gauss' or two for 'ring' kernel, see details.
- **with_cov**: logical. If TRUE not only spatial covariance matrices but also the sample covariances matrices for each sub-domain are considered in the joint diagonalization procedure. Default is TRUE.
- **lcov**: a string indicating which type of local covariance matrix to use. Either 'lcoov' (default), 'ldiff' or 'lcoov_norm'. See sbss_asymp for details on the latter option.
- **ordered**: logical. If TRUE the entries of the latent field are ordered by the sum of squared pseudo-eigenvalues of the diagonalized sub-domain (local) covariance matrices. Default is TRUE.
- **...**: further arguments for the fast real joint diagonalization algorithm that jointly diagonalizes the sub-domain covariance matrices. See details and frjd.

### Details

This function assumes that the random field \( x \) is formed by

\[
x(t) = As(t) + b,
\]
where $A$ is the deterministic $p \times p$ mixing matrix, $b$ is the $p$-dimensional location vector, $x$ is the observable $p$-variate random field given by the argument $x$, $t$ are the spatial locations given by the argument coords and $s$ is the latent $p$-variate random field assumed to consist of uncorrelated entries that have zero mean but non-constant (spatial) second order dependence. This function aims to recover $s$ by

$$W(x(t) - \bar{x}),$$

where $W$ is the $p \times p$ unmixing matrix and $\bar{x}$ is the sample mean. The function does this by splitting the given spatial domain into $n_{\text{block}}^2$ equally sized rectangular sub-domains and jointly diagonalizing the corresponding spatial covariance matrices for all sub-domains. If the argument with_cov equals TRUE (default) then additionally also the sample covariance matrices for each sub-domain are included in the joint diagonalization procedure.

The arguments kernel_type, kernel_parameters and lcov determine which spatial kernel functions and which type of local covariance matrices are used for each sub-domain. The usage is equal to the function sbss.

Alternatively the domain subdivision can be defined by providing lists of length $K$ for the arguments $x$ and coords where the first list entries correspond to the values and coordinates of the first sub-domain and the second entries to the values and coordinates of the second sub-domain, etc.. The argument $n_{\text{block}}$ might be 'x' or 'y' indicating a split across the x or y coordinates similar as done by the function snss_sd.

snss_sjd jointly diagonalizes the covariance matrices for each sub-domain with the function frjd. ... provides arguments for frjd, useful arguments might be:

- eps: tolerance for convergence.
- maxiter: maximum number of iterations.

Value

Similarly as sbss the function snss_jd returns a list of class 'snss' and 'sbss' with the following entries:

- s: object of class(x) containing the estimated source random field.
- coords: coordinates of the observations. Only given if $x$ is a matrix or list.
- w: estimated unmixing matrix.
- w_inv: inverse of the estimated unmixing matrix.
- d: matrix of stacked (jointly) diagonalized sub-domain covariance and/or local covariance matrices.
- x_mu: columnmeans of $x$.
- cov_inv_sqrt: square root of the inverse sample covariance matrix of $x$.

References


See Also

sbss, sp, sf
Examples

# simulate coordinates
n <- 1000
coords <- runif(n * 2) * 20
dim(coords) <- c(n, 2)

# simulate random field
field_1 <- rnorm(n)
field_2 <- 2 * sin(pi / 20 * coords[, 1]) * rnorm(n)
field_3 <- rnorm(n) * (coords[, 1] < 10) + rnorm(n, 0, 3) * (coords[, 1] >= 10)

latent_field <- cbind(field_1, field_2, field_3)
mixing_matrix <- matrix(rnorm(9), 3, 3)
observed_field <- latent_field

observed_field_sp <- sp::SpatialPointsDataFrame(coords = coords,
                                             data = data.frame(observed_field))
sp::spplot(observed_field_sp, colorkey = TRUE, as.table = TRUE, cex = 1)

# apply snss_sjd with 4 sub-domains
# one ring kernel per sub-domain
# without covariances
res_4_ball <- snss_sjd(observed_field, coords, n_block = 2,
                        kernel_type = 'ball', kernel_parameters = c(0, 2),
                        with_cov = TRUE)
JADE::MD(W.hat = coef(res_4_ball), A = mixing_matrix)

# apply snss_sjd with split across y
# one ring kernel per sub-domain
# without covariances
# should not work as field does not show spatial dependence
res_4_ring <- snss_sjd(observed_field, coords, n_block = 'y',
                       kernel_type = 'ring', kernel_parameters = c(0, 2),
                       with_cov = FALSE)
JADE::MD(W.hat = coef(res_4_ring), A = mixing_matrix)

# print object
print(res_4_ball)

# plot latent field
plot(res_4_ball, colorkey = TRUE, as.table = TRUE, cex = 1)

# predict latent fields on grid
predict(res_4_ball, colorkey = TRUE, as.table = TRUE, cex = 1)

# unmixing matrix
w_unmix <- coef(res_4_ball)

# apply snss_jd with SpatialPointsDataFrame object
res_4_ball_sp <- snss_sjd(observed_field_sp, n_block = 2,
                         kernel_type = 'ball', kernel_parameters = c(0, 2),
                         with_cov = TRUE)
# apply with list arguments
# first axis split by 5
# second axis split by 10
# results in 4 sub-domains
flag_x <- coords[, , 1] < 5
flag_y <- coords[, , 2] < 10
coords_list <- list(coords[flag_x & flag_y, ],
                     coords[!flag_x & flag_y, ],
                     coords[flag_x & !flag_y, ],
                     coords[!flag_x & !flag_y, ])
field_list <- list(observed_field[flag_x & flag_y, ],
                   observed_field[!flag_x & flag_y, ],
                   observed_field[flag_x & !flag_y, ],
                   observed_field[!flag_x & !flag_y, ]) plot(coords, col = 1)
points(coords_list[[2]], col = 2)
points(coords_list[[3]], col = 3)
points(coords_list[[4]], col = 4)
res_list <- snss_sjd(x = field_list,
                     coords = coords_list,
                     kernel_type = 'ring', kernel_parameters = c(0, 2))
plot(res_list, colorkey = TRUE, as.table = TRUE, cex = 1)
JADE::MD(W.hat = coef(res_list), A = mixing_matrix)

---

**spatial_kernel_matrix**  
*Computation of Spatial Kernel Matrices*

**Description**

`spatial_kernel_matrix` computes spatial kernel matrices for a given kernel function with its parameters and a set of coordinates.

**Usage**

`spatial_kernel_matrix(coords, kernel_type = c('ring', 'ball', 'gauss'),
                       kernel_parameters)`

**Arguments**

- `coords` a numeric matrix of dimension c(n,2) where each row represents the coordinates of a point in the spatial domain.
- `kernel_type` a character string indicating which kernel function to use. Either 'ring' (default), 'ball' or 'gauss'.
- `kernel_parameters` a numeric vector that gives the parameters for the kernel function. At least length of one for 'ball' and 'gauss' or two for 'ring' kernel, see details.
Details

Two versions of local covariance matrices can be defined:

- 'lcov':
  \[ LCov(f) = \frac{1}{n} \sum_{i,j} f(d_{i,j})(x(s_i) - \bar{x})(x(s_j) - \bar{x})' \]

- 'ldiff':
  \[ LDiff(f) = \frac{1}{n} \sum_{i,j} f(d_{i,j})(x(s_i) - x(s_j))(x(s_i) - x(s_j))' \]

- 'lcov_norm':
  \[ LCov^*(f) = \frac{1}{n F_{f,n}^{1/2}} \sum_{i,j} f(d_{i,j})(x(s_i) - \bar{x})(x(s_j) - \bar{x})' \]

with

\[ F_{f,n} = \frac{1}{n} \sum_{i,j} f^2(d_{i,j}). \]

Where \( d_{i,j} \geq 0 \) correspond to the pairwise distances between coordinates, \( x(s_i) \) are the \( p \) random field values at location \( s_i \), \( \bar{x} \) is the sample mean vector, and the kernel function \( f(d) \) determines the locality. The function \( \text{spatial_kernel_matrix} \) computes a list of \( c(n,n) \) matrices where each entry of these matrices correspond to the spatial kernel function evaluated at the distance between two points, mathematically the entry \( ij \) of each kernel matrix is \( f(d_{i,j}) \). The following kernel functions are implemented and chosen with the argument \( \text{kernel_type} \):

- 'ring': parameters are inner radius \( r_i \) and outer radius \( r_o \), with \( r_i < r_o \), and \( r_i, r_o \geq 0 \):
  \[ f(d; r_i, r_o) = I(r_i < d \leq r_o) \]

- 'ball': parameter is the radius \( r \), with \( r \geq 0 \):
  \[ f(d; r) = I(d \leq r) \]

- 'gauss': Gaussian function where 95% of the mass is inside the parameter \( r \), with \( r \geq 0 \):
  \[ f(d; r) = \exp(-0.5(\Phi^{-1}(0.95)d/r)^2) \]

The argument \( \text{kernel_type} \) determines the used kernel function as presented above, the argument \( \text{kernel_parameters} \) gives the corresponding parameters for the kernel function. Specifically, if \( \text{kernel_type} \) equals 'ball' or 'gauss' then \( \text{kernel_parameters} \) is a numeric vector where each entry corresponds to one parameter. Hence, \( \text{length(} \text{kernel_parameters} \text{)} \) spatial kernel matrices of type \( \text{kernel_type} \) are computed. Whereas, if \( \text{kernel_type} \) equals 'ring', then \( \text{kernel_parameters} \) must be a numeric vector of even length where subsequently the inner and outer radii must be given (informally: \( c(r_{i1}, r_{o1}, r_{i2}, r_{o2}, ...) \)). In that case \( \text{length(} \text{kernel_parameters} \text{)} / 2 \) spatial kernel matrices of type 'ring' are computed.

The output of this function can be used with the function \( \text{sbss} \) to avoid unnecessary computation of kernel matrices when \( \text{sbss} \) is called multiple times with the same coordinate/kernel function setting. Additionally, the output can be used with the function \( \text{local_covariance_matrix} \) to actually compute local covariance matrices as defined above based on a given set of spatial kernel matrices.
Value

 spatial_kernel_matrix returns a list with length of length(kernel_parameters) (for 'ball' and 'gauss' kernel functions) or length(kernel_parameters) / 2 (for 'ring' kernel function) containing numeric matrices of dimension c(n,n) corresponding to the spatial kernel matrices.

References


See Also

sbss, local_covariance_matrix

Examples

# simulate a set of coordinates
coords <- rnorm(100 * 2)
dim(coords) <- c(100, 2)

# computing two ring kernel matrices
kernel_params_ring <- c(0, 0.5, 0.5, 2)
ring_kernel_list <-
 spatial_kernel_matrix(coords, 'ring', kernel_params_ring)

# computing three ball kernel matrices
kernel_params_ball <- c(0.5, 1, 2)
ball_kernel_list <-
 spatial_kernel_matrix(coords, 'ball', kernel_params_ball)

# computing three gauss kernel matrices
kernel_params_gauss <- c(0.5, 1, 2)
gauss_kernel_list <-
 spatial_kernel_matrix(coords, 'gauss', kernel_params_gauss)

---

white_data

Different Approaches of Data Whitening

Description

white_data whites the data with respect to the sample covariance matrix, or different spatial scatter matrices.
Usage

white_data(x, whitening = c("standard", "rob", "hr"),
   lcov = c('lcov', 'ldiff', 'lcov_norm'),
   kernel_mat = numeric(0))

Arguments

x a numeric matrix of dimension c(n, p) where the p columns correspond to the entries of the random field and the n rows are the observations.
whitening a string indicating the whitening method. If 'standard' then the whitening is carried out with respect to sample covariance matrix, if 'rob' then the first spatial scatter matrix is used instead of sample the covariance matrix and if 'hr' then the Hettmansperger-Randles location and scatter estimates are used for whitening. See details for more. Default is 'standard'.
lcov a string indicating which type of local covariance matrix is used for whitening, when the whitening method 'rob' is used. Either 'lcov' (default) or 'ldiff'.
kernel_mat a spatial kernel matrix with dimension c(n,n), see details. Usually computed by the function spatial_kernel_matrix.

Details

The inverse square root of a positive definite matrix $M(x)$ with eigenvalue decomposition $UDU'$ is defined as $M(x)^{-1/2} = UD^{-1/2}U'$. white_data whitens the data by $M(x)^{-1/2}(x - T(x))$ where $T(x)$ is a location functional of $x$ and the matrix $M(x)$ is a scatter functional. If the argument whitening is 'standard', $M(x)$ is the sample covariance matrix and $T(x)$ is a vector of column means of $x$. If the argument whitening is 'hr', the Hettmansperger-Randles location and scatter estimates (Hettmansperger & Randles, 2002) are used as location functional $T(x)$ and scatter functional $M(x)$. The Hettmansperger-Randles location and scatter estimates are robust variants of sample mean and covariance matrices, that are used for whitening in robsbss. If the argument whitening is 'rob', the argument lcov determines the scatter functional $M(x)$ to be one of the following local scatter matrices:

- 'lcov':
  $$LCov(f) = \frac{1}{n} \sum_{i,j} f(d_{i,j})(x(s_i) - \bar{x})(x(s_j) - \bar{x})',$$

- 'ldiff':
  $$LDiff(f) = \frac{1}{n} \sum_{i,j} f(d_{i,j})(x(s_i) - x(s_j))(x(s_i) - x(s_j))',$$

- 'lcov_norm':
  $$LCov^*(f) = \frac{1}{(nF_{f,n}^{1/2})} \sum_{i,j} f(d_{i,j})(x(s_i) - \bar{x})(x(s_j) - \bar{x})',$$

with

$$F_{f,n} = \frac{1}{n} \sum_{i,j} f^2(d_{i,j}),$$
where \( d_{i,j} \geq 0 \) correspond to the pairwise distances between coordinates, \( x(s_i) \) are the \( p \) random field values at location \( s_i \), \( \bar{x} \) is the sample mean vector, and the kernel function \( f(d) \) determines the locality. The choice ‘\texttt{lcov_norm}’ is useful when testing for the actual signal dimension of the latent field, see \texttt{sbss_asym} and \texttt{sbss_boot}. See also \texttt{sbss} for details.

Note that \( LCov(f) \) are usually not positive definite, therefore in that case the matrix cannot be inverted and an error is produced. Whitening with \( LCov(f) \) matrices might be favorable in the presence of spatially uncorrelated noise, and whitening with \( LDiff(f) \) might be favorable when a non-constant smooth drift is present in the data.

The argument \texttt{kernel_mat} is a matrix of dimension \( c(n,n) \) where each entry corresponds to the spatial kernel function evaluated at the distance between two sample locations, mathematically the entry \( ij \) of each kernel matrix is \( f(d_{i,j}) \). This matrix is usually computed with the function \texttt{spatial_kernel_matrix}.

**Value**

\texttt{white_data} returns a list with the following entries:

- \texttt{mu} a numeric vector of length \( ncol(x) \) containing the column means of the data matrix \( x \).
- \texttt{x_0} a numeric matrix of dimension \( c(n, p) \) containing the columns centered data of \( x \).
- \texttt{x_w} a numeric matrix of dimension \( c(n, p) \) containing the whitened data of \( x \).
- \texttt{s} a numeric matrix of dimension \( c(p, p) \) which is the scatter matrix \( M \).
- \texttt{s_inv_sqrt} a numeric matrix of dimension \( c(p, p) \) which equals the inverse square root of the scatter matrix \( M \) used for whitening.
- \texttt{s_sqrt} a numeric matrix of dimension \( c(p, p) \) which equals the square root of the scatter matrix \( M \).

**References**


**See Also**

\texttt{sbss, spatial_kernel_matrix}

**Examples**

```r
# simulate coordinates
coords <- runif(1000 * 2) * 20
dim(coords) <- c(1000, 2)
```
coords_df <- as.data.frame(coords)
names(coords_df) <- c("x", "y")
# simulate random field
if (!requireNamespace("gstat", quietly = TRUE)) {
  message("Please install the package gstat to run the example code."
} else {
  library(gstat)
  model_1 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                    model = vgm(psill = 0.025, range = 1, model = 'Exp'), nmax = 20)
  model_2 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                    model = vgm(psill = 0.025, range = 1, kappa = 2, model = 'Mat'),
                           nmax = 20)
  model_3 <- gstat(formula = z ~ 1, locations = ~ x + y, dummy = TRUE, beta = 0,
                    model = vgm(psill = 0.025, range = 1, model = 'Gau'), nmax = 20)
  field_1 <- predict(model_1, newdata = coords_df, nsim = 1)$sim1
  field_2 <- predict(model_2, newdata = coords_df, nsim = 1)$sim1
  field_3 <- predict(model_3, newdata = coords_df, nsim = 1)$sim1
  field <- cbind(field_1, field_2, field_3)
  X <- as.matrix(field)

  # white the data with the usual sample covariance
  x_w_1 <- white_data(X)

  # white the data with a ldiff matrix and ring kernel
  kernel_params_ring <- c(0, 1)
  ring_kernel_list <-
    spatial_kernel_matrix(coords, 'ring', kernel_params_ring)
  x_w_2 <- white_data(field, whitening = 'rob',
                      lcov = 'ldiff', kernel_mat = ring_kernel_list[[1]])

  # Generate 5 % of global outliers to data
  field_cont <- gen_glob_outl(field)[,1:3]
  X <- as.matrix(field_cont)
  # white the data using Hettmansperger-Randles location and scatter estimates
  x_w_3 <- white_data(X, whitening = 'hr')
}
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