Package ‘convoSPAT’

January 16, 2021

Type Package
Title Convolution-Based Nonstationary Spatial Modeling
Version 1.2.7
Date 2021-01-15
Description Fits convolution-based nonstationary
Gaussian process models to point-referenced spatial data. The nonstationary
covariance function allows the user to specify the underlying correlation
structure and which spatial dependence parameters should be allowed to
vary over space: the anisotropy, nugget variance, and process variance.
The parameters are estimated via maximum likelihood, using a local
likelihood approach. Also provided are functions to fit stationary spatial
models for comparison, calculate the Kriging predictor and standard errors,
and create various plots to visualize nonstationarity.

Depends R (>= 3.1.2)
License MIT + file LICENSE
LazyData TRUE
Imports stats, graphics, ellipse, fields, MASS, plotrix, StatMatch
URL http://github.com/markdrisser/convoSPAT
RoxygenNote 7.1.0

R topics documented:

Aniso_fit .......................................................... 2
cov.spatial .......................................................... 5
evaluate_CV ......................................................... 5
Aniso_fit

Fit the stationary spatial model

Description

Aniso_fit estimates the parameters of the stationary spatial model. Required inputs are the observed data and locations. Optional inputs include the covariance model (exponential is the default).

Usage

Aniso_fit(
  sp.SPDF = NULL,
  coords = NULL,
  data = NULL,
  cov.model = "exponential",
  mean.model = data ~ 1,
  fixed.nugg2.var = NULL,
  method = "reml",
  fix.tausq = FALSE,
  tausq = 0,
  fix.kappa = FALSE,
  kappa = 0.5,
local.pars.LB = NULL,
local.pars.UB = NULL,
local.ini.pars = NULL
)

Arguments

sp.SPDF A "SpatialPointsDataFrame" object, which contains the spatial coordinates and additional attribute variables corresponding to the spatial coordinates

cov.model A string specifying the model for the correlation function; defaults to "exponential". Options available in this package are: "exponential", "matern", or "gaussian".

mean.model An object of class formula, specifying the mean model to be used. Defaults to an intercept only.

fixed.nugg2.var Optional; describes the variance/covariance for a fixed (second) nugget term (represents a known error term). Either a vector of length N containing a station-specific variances (implying independent error) or an N x N covariance matrix (implying dependent error). Defaults to zero.

method Indicates the estimation method, either maximum likelihood ("ml") or restricted maximum likelihood ("reml").

fix.tausq Logical; indicates whether the default nugget term (tau^2) should be fixed (TRUE) or estimated (FALSE). Defaults to FALSE.

tausq Scalar; fixed value for the nugget variance (when fix.tausq = TRUE).

fix.kappa Logical; indicates if the kappa parameter should be fixed (TRUE) or estimated (FALSE). Defaults to FALSE (only valid for cov.model = "matern" and cov.model = "cauchy").

kappa Scalar; value of the kappa parameter. Only used if fix.kappa = TRUE.

local.pars.LB, local.pars.UB Optional vectors of lower and upper bounds, respectively, used by the "L-BFGS-B" method option in the optim function for the local parameter estimation. Each vector must be of length five, containing values for lam1, lam2, tausq, sigmasq, and nu. Default for local.pars.LB is rep(1e-05,5); default for local.pars.UB is c(max.distance/2,max.distance/2,4*resid.var,4*resid.var,100), where max.distance is the maximum interpoint distance of the observed data and resid.var is the residual variance from using lm with mean.model.

local.ini.pars Optional vector of initial values used by the "L-BFGS-B" method option in the optim function for the local parameter estimation. The vector must be of length five, containing values for lam1, lam2, tausq, sigmasq, and nu. Defaults to c(max.distance/10,max.distance/10,0.1*resid.var,0.9*resid.var,1), where max.distance is the maximum interpoint distance of the observed data and resid.var is the residual variance from using lm with mean.model.
Value

A list with the following components:

- **MLEs.save**: Table of local maximum likelihood estimates for each mixture component location.
- **data**: Observed data values.
- **beta.GLS**: Vector of generalized least squares estimates of beta, the mean coefficients.
- **beta.cov**: Covariance matrix of the generalized least squares estimate of beta.
- **Mean.coefs**: "Regression table" for the mean coefficient estimates, listing the estimate, standard error, and t-value.
- **Cov.mat**: Estimated covariance matrix (N.obs x N.obs) using all relevant parameter estimates.
- **Cov.mat.chol**: Cholesky of Cov.mat (i.e., chol(Cov.mat)), the estimated covariance matrix (N.obs x N.obs).
- **aniso.pars**: Vector of MLEs for the anisotropy parameters lam1, lam2, eta.
- **aniso.mat**: 2 x 2 anisotropy matrix, calculated from aniso.pars.
- **tausq.est**: Scalar maximum likelihood estimate of tausq (nugget variance).
- **sigmasq.est**: Scalar maximum likelihood estimate of sigmasq (process variance).
- **kappa.MLE**: Scalar maximum likelihood estimate for kappa (when applicable).
- **fixed.nugg2.var**: N x N matrix with the fixed variance/covariance for the second (measurement error) nugget term (defaults to zero).
- **cov.model**: String; the correlation model used for estimation.
- **coords**: N x 2 matrix of observation locations.
- **global.loglik**: Scalar value of the maximized likelihood from the global optimization (if available).
- **Xmat**: Design matrix, obtained from using lm with mean.model.
- **fix.kappa**: Logical, indicating if kappa was fixed (TRUE) or estimated (FALSE).
- **kappa**: Scalar; fixed value of kappa.

Examples

```r
## Not run:
# Using iid standard Gaussian data
aniso.fit <- Aniso_fit( coords = cbind(runif(100), runif(100)),
                       data = rnorm(100) )
## End(Not run)
```
cov_spatial

Calculate spatial covariance.

Description
This function replaces the geoR function cov.spatial, which is now defunct. Options available in this package are: "exponential", "matern", and "gaussian".

Usage

cov_spatial(
  Dist.mat,
  cov.model = "exponential",
  cov.pars = c(1, 1),
  kappa = 0.5
)

Arguments
Dist.mat A matrix of scaled distances.
cov.model A string specifying the model for the correlation function; defaults to "exponential". Options available in this package are: "exponential", "matern", and "gaussian".
cov.pars Fixed values; not used in the function.
kappa Scalar; value of the smoothness parameter.

Value
This function returns a correlation matrix.

Examples

Distmat <- as.matrix(dist(matrix(runif(20), ncol = 2), diag = TRUE, upper = TRUE))
C <- cov_spatial( Dist.mat = Distmat )

evaluate.CV

Evaluation criteria

Description
Calculate three evaluation criteria – continuous rank probability score (CRPS), prediction mean square deviation ratio (pMSDR), and mean squared prediction error (MSPE) – comparing hold-out data and predictions.
Usage

\texttt{evaluate.CV(holdout.data, pred.mean, pred.SDs)}

Arguments

\begin{itemize}
  \item \texttt{holdout.data} Observed/true data that has been held out for model comparison.
  \item \texttt{pred.mean} Predicted mean values corresponding to the hold-out locations.
  \item \texttt{pred.SDs} Predicted standard errors corresponding to the hold-out locations.
\end{itemize}

Value

A list with the following components:

\begin{itemize}
  \item \texttt{CRPS} The CRPS averaged over all hold-out locations.
  \item \texttt{MSPE} The mean squared prediction error.
  \item \texttt{pMSDR} The prediction mean square deviation ratio.
\end{itemize}

Examples

\begin{verbatim}
## Not run:
evaluate.CV( holdout.data = simdata$sim.data[holdout.index],
pred.mean = pred.NS$pred.means, pred.SDs = pred.NS$pred.SDs )
## End(Not run)
\end{verbatim}

\textbf{f.mc.kernels} \hspace{1cm} \textit{Calculate mixture component kernel matrices.}

Description

\texttt{f.mc.kernels} calculates spatially-varying mixture component kernels using generalized linear models for each of the eigenvalues (lam1 and lam2) and the angle of rotation (eta).

Usage

\begin{verbatim}
f.mc.kernels(
  y.min = 0,
  y.max = 5,
  x.min = 0,
  x.max = 5,
  N.mc = 3^2,
  lam1.coef = c(-1.3, 0.5, -0.6),
  lam2.coef = c(-1.4, -0.1, 0.2),
  logit.eta.coef = c(0, -0.15, 0.15)
)
\end{verbatim}
kernel_cov

Arguments

y.min Lower bound for the y-coordinate axis.
y.max Upper bound for the y-coordinate axis.
x.min Lower bound for the y-coordinate axis.
x.max Upper bound for the y-coordinate axis.
N.mc Number of mixture component locations.
lam1.coef Log-linear regression coefficients for lam1; the coefficients correspond to the intercept, longitude, and latitude.
lam2.coef Log-linear regression coefficients for lam2; the coefficients correspond to the intercept, longitude, and latitude.
logit.eta.coef Scaled logit regression coefficients for eta; the coefficients correspond to the intercept, longitude, and latitude.

Value

A list with the following components:

mc.locations A N.mc x 2 matrix of the mixture component locations.
mc.kernels A N.mc x 2 x 2 array of kernel matrices corresponding to each of the mixture component locations.

Examples

f_mc_kernels(y.min = 0, y.max = 5, x.min = 0,
x.max = 5, N.mc = 3^2, lam1.coef = c(-1.3, 0.5, -0.6),
lam2.coef = c(-1.4, -0.1, 0.2), logit.eta.coef = c(0, -0.15, 0.15) )

kernel_cov

Calculate a kernel covariance matrix.

Description

kernel_cov calculates a 2 x 2 matrix based on the eigendecomposition components (two eigenvalues and angle of rotation).

Usage

kernel_cov(params)

Arguments

params A vector of three parameters, corresponding to (lam1, lam2, eta). The eigenvalues (lam1 and lam2) must be positive.
Value

A 2 x 2 kernel covariance matrix.

Examples

kernel_cov(c(1, 2, pi/3))

Description

This function generates another function to be used within `optim` to obtain maximum likelihood estimates of global variance parameters tau^2, sigma^2 with a fixed correlation matrix (smoothness is fixed).

Usage

```r
make_global_loglik1(data, Xmat, Corr, nugg2.var)
```

Arguments

- `data` A vector or matrix of data to use in the likelihood calculation.
- `Xmat` The design matrix for the mean model.
- `Corr` The correlation matrix.
- `nugg2.var` Fixed values for the covariance of the second nugget term.

Value

This function returns another function for use in `optim`.

Examples

```r
## Not run:
make_global_loglik1( data, Xmat, Corr, nugg2.var )
## End(Not run)
```
make_global_loglik1_kappa

Constructor functions for global parameter estimation.

Description

This function generates another function to be used within optim to obtain maximum likelihood estimates of global variance parameters tausq, sigmasq, and nu.

Usage

make_global_loglik1_kappa(data, Xmat, cov.model, Scalemat, Distmat, nugg2.var)

Arguments

data: A vector or matrix of data to use in the likelihood calculation.
Xmat: The design matrix for the mean model.
cov.model: String; the covariance model.
Scalemat: Matrix; contains the scaling quantities from the covariance function.
Distmat: Matrix; contains the scaled distances.
nugg2.var: Fixed values for the covariance of the second nugget term.

Value

This function returns another function for use in optim.

Examples

### Not run:
make_global_loglik1_kappa( data, Xmat, cov.model, Scalemat, Distmat, nugg2.var )

### End(Not run)

make_global_loglik2 Constructor functions for global parameter estimation.

Description

This function generates another function to be used within optim to obtain maximum likelihood estimates of global variance parameter sigmasq with a fixed correlation matrix (smoothness is fixed). The nugget variance is taken to be spatially-varying.
Usage

make_global_loglik2(data, Xmat, Corr, obs.nuggets, nugg2.var)

Arguments

data A vector or matrix of data to use in the likelihood calculation.
Xmat The design matrix for the mean model.
Corr The correlation matrix.
obs.nuggets A vector containing the spatially-varying nuggets corresponding to each data location.
nugg2.var Fixed values for the covariance of the second nugget term.

Value

This function returns another function for use in optim.

Examples

## Not run:
make_global_loglik2( data, Xmat, Corr, obs.nuggets, nugg2.var )
## End(Not run)
make_global_loglik3

Arguments

- **data**: A vector or matrix of data to use in the likelihood calculation.
- **Xmat**: The design matrix for the mean model.
- **cov.model**: String; the covariance model.
- **Scalemat**: Matrix; contains the scaling quantities from the covariance function.
- **Distmat**: Matrix; contains the scaled distances.
- **obs.nuggets**: A vector containing the spatially-varying nuggets corresponding to each data location.
- **nugg2.var**: Fixed values for the covariance of the second nugget term.

Value

This function returns another function for use in `optim`.

Examples

```r
## Not run:
make_global_loglik2_kappa( data, Xmat, cov.model, Scalemat, Distmat, obs.nuggets, nugg2.var )
## End(Not run)
```

make_global_loglik3  Constructor functions for global parameter estimation.

Description

This function generates another function to be used within `optim` to obtain maximum likelihood estimates of global variance parameter tau_sq with a fixed correlation matrix (smoothness is fixed). The process variance is taken to be spatially-varying.

Usage

```r
make_global_loglik3(data, Xmat, Corr, obs.variance, nugg2.var)
```

Arguments

- **data**: A vector or matrix of data to use in the likelihood calculation.
- **Xmat**: The design matrix for the mean model.
- **Corr**: The correlation matrix matrix.
- **obs.variance**: A vector containing the spatially-varying variance corresponding to each data location.
- **nugg2.var**: Fixed values for the covariance of the second nugget term.
This function returns another function for use in \texttt{optim}.

Examples

```r
## Not run:
make_global_loglik3( data, Xmat, Corr, obs.variance, nugg2.var )
## End(Not run)
```

make\_global\_loglik3\_kappa

\textit{Constructor functions for global parameter estimation.}

**Description**

This function generates another function to be used within \texttt{optim} to obtain maximum likelihood estimates of global variance parameters tausq and nu. The process variance is taken to be spatially-varying.

**Usage**

```r
make_global_loglik3_kappa(  
  data,  
  Xmat,  
  cov.model,  
  Scalemat,  
  Distmat,  
  obs.variance,  
  nugg2.var
)
```

**Arguments**

- **data**: A vector or matrix of data to use in the likelihood calculation.
- **Xmat**: The design matrix for the mean model.
- **cov.model**: String; the covariance model.
- **Scalemat**: Matrix; contains the scaling quantities from the covariance function.
- **Distmat**: Matrix; contains the scaled distances.
- **obs.variance**: A vector containing the spatially-varying variance corresponding to each data location.
- **nugg2.var**: Fixed values for the covariance of the second nugget term.
Value

This function returns another function for use in optim.

Examples

```r
## Not run:
make_global_loglik3_kappa( data, Xmat, cov.model, Scalemat, Distmat, obs.variance, nugg2.var )
## End(Not run)
```

Description

This function generates another function to be used within optim to obtain maximum likelihood estimates of global variance parameters \( \nu \). The process variance and nugget variance are taken to be spatially-varying.

Usage

```r
make_global_loglik4_kappa(
  data,
  Xmat,
  cov.model,
  Scalemat,
  Distmat,
  obs.variance,
  obs.nuggets,
  nugg2.var
)
```

Arguments

- `data` A vector or matrix of data to use in the likelihood calculation.
- `Xmat` The design matrix for the mean model.
- `cov.model` String; the covariance model.
- `Scalemat` Matrix; contains the scaling quantities from the covariance function.
- `Distmat` Matrix; contains the scaled distances.
- `obs.variance` A vector containing the spatially-varying variance corresponding to each data location.
- `obs.nuggets` A vector containing the spatially-varying nuggets corresponding to each data location.
- `nugg2.var` Fixed values for the covariance of the second nugget term.
Value

This function returns another function for use in optim.

Examples

## Not run:
make_global_loglik4_kappa( data, Xmat, cov.model, Scalemat, Distmat, 
obs.variance, obs.nuggets, nugg2.var )

## End(Not run)

make_local_lik Constructor functions for local parameter estimation.

Description

This function generates another function to be used within optim to obtain maximum likelihood estimates of covariance (and possibly mean) parameters. The function includes options for (1) maximum likelihood ("ml") vs. restricted maximum likelihood ("reml"), (2) smoothness (kappa): models without smoothness vs. estimating the smoothness vs. using fixed smoothness, (3) locally isotropic vs. locally anisotropic, and (4) fixed nugget variance (tausq): fixed vs. estimated.

Usage

make_local_lik(
  locations,
  cov.model,
  data,
  Xmat,
  nugg2.var = matrix(0, nrow(locations), nrow(locations)),
  tausq = 0,
  kappa = 0.5,
  fixed = rep(FALSE, 6),
  method = "reml",
  local.aniso = TRUE,
  fix.tausq = FALSE,
  fix.kappa = FALSE
)

Arguments

  locations      A matrix of locations.
  cov.model     String; the covariance model.
  data          A vector or matrix of data to use in the likelihood calculation.
  Xmat          The design matrix for the mean model.
**mc_N**

`mc_N` calculates the number of observations (sample size) that fall within a certain fit radius for each mixture component location.

### Description

`mc_N` calculates the number of observations (sample size) that fall within a certain fit radius for each mixture component location.

### Usage

```r
mc_N(coords, mc.locations, fit.radius)
```

### Arguments

- **coords**: A matrix of observation locations.
- **mc.locations**: A matrix of the mixture component locations to use in the model fitting.
- **fit.radius**: Scalar; defines the fitting radius for local likelihood estimation.
Value

A vector `mc.N.fit`, which summarizes the number of observation locations in `coords` that fall within the fit radius for each mixture component location.

Examples

```r
## Not run:
mc_N(coords = simdata$sim.locations, mc.locations = simdata$mc.locations,
     fit.radius = 1 )

## End(Not run)
```

Description

`NSconvo_fit` estimates the parameters of the nonstationary convolution-based spatial model. Required inputs are the observed data and locations. Optional inputs include mixture component locations (if not provided, the number of mixture component locations are required), the fit radius, the covariance model (exponential is the default), and whether or not the nugget and process variance will be spatially-varying.

Usage

```r
NSconvo_fit(sp.SPDF = NULL, coords = NULL, data = NULL, cov.model = "exponential",
             mean.model = data ~ 1, mc.locations = NULL, N.mc = NULL,
             lambda.w = NULL, fixed.nugg2.var = NULL, mean.model.df = NULL,
             mc.kernels = NULL, fit.radius = NULL, ns.nugget = FALSE,
             ns.variance = FALSE, ns.mean = FALSE, local.aniso = TRUE,
             fix.tausq = FALSE, tausq = 0, fix.kappa = FALSE, kappa = 0.5,
```

method = "reml",
print.progress = TRUE,
local.pars.LB = NULL,
local.pars.UB = NULL,
global.pars.LB = NULL,
global.pars.UB = NULL,
local.ini.pars = NULL,
global.ini.pars = NULL
)

Arguments

sp.SPDF A "SpatialPointsDataFrame" object, which contains the spatial coordinates and additional attribute variables corresponding to the spatial coordinates.

cords An N x 2 matrix where each row has the two-dimensional coordinates of the N data locations.

data A vector or matrix with N rows, containing the data values. Inputting a vector corresponds to a single replicate of data, while inputting a matrix corresponds to replicates. In the case of replicates, the model assumes the replicates are independent and identically distributed.

cov.model A string specifying the model for the correlation function; defaults to "exponential". Options available in this package are: "exponential", "matern", and "gaussian".

mean.model An object of class formula, specifying the mean model to be used. Defaults to an intercept only.

mc.locations Optional; matrix of mixture component locations.

N.mc Optional; if mc.locations is not specified, the function will create a rectangular grid of size N.mc over the spatial domain.

lambda.w Scalar; tuning parameter for the weight function. Defaults to be the square of one-half of the minimum distance between mixture component locations.

fixed.nugg2.var Optional; describes the variance/covariance for a fixed (second) nugget term (represents a known error term). Either a vector of length N containing a station-specific variances (implying independent error) or an NxN covariance matrix (implying dependent error). Defaults to zero.

mean.model.df Optional data frame; refers to the variables used in mean.model. Important when using categorical variables in mean.model, as a subset of the full design matrix will likely be rank deficient. Specifying mean.model.df allows NSconvo_fit to calculate a design matrix specific to the points used to fit each local model.

mc.kernels Optional specification of mixture component kernel matrices (based on expert opinion, etc.).

fit.radius Scalar; specifies the fit radius or neighborhood size for the local likelihood estimation.

ns.nugget Logical; indicates if the nugget variance (tausq) should be spatially-varying (TRUE) or constant (FALSE).
**ns.variance**
Logical; indicates if the process variance (sigmasq) should be spatially-varying (TRUE) or constant (FALSE).

**ns.mean**
Logical; indicates if the mean coefficients (beta) should be spatially-varying (TRUE) or constant (FALSE).

**local.aniso**
Logical; indicates if the local covariance should be anisotropic (TRUE) or isotropic (FALSE). Defaults to TRUE. In the case of a locally isotropic model, the bounds and initial values for lam will default to the first element of local.pars.LB, local.pars.UB, and local.ini.pars (while still required, the second and third elements of these vectors will be ignored.)

**fix.tausq**
Logical; indicates whether the default nugget term (tau^2) should be fixed (TRUE) or estimated (FALSE). Defaults to FALSE.

**tausq**
Scalar; fixed value for the nugget variance (when fix.tausq = TRUE).

**fix.kappa**
Logical; indicates if the kappa parameter should be fixed (TRUE) or estimated (FALSE). Defaults to FALSE (only valid for cov.model = "matern" and cov.model = "cauchy").

**kappa**
Scalar; value of the kappa parameter. Only used if fix.kappa = TRUE.

**method**
Indicates the estimation method, either maximum likelihood ("ml") or restricted maximum likelihood ("reml").

**print.progress**
Logical; if TRUE, text indicating the progress of local model fitting in real time.

**local.pars.LB, local.pars.UB**
Optional vectors of lower and upper bounds, respectively, used by the "L-BFGS-B" method option in the optim function for the local parameter estimation. Each vector must be of length five, containing values for lam1, lam2, tausq, sigmasq, and nu. Default for local.pars.LB is rep(1e-05,5); default for local.pars.UB is c(max.distance/2,max.distance/2,4*resid.var,4*resid.var,100), where max.distance is the maximum interpoint distance of the observed data and resid.var is the residual variance from using lm with mean.model.

**global.pars.LB, global.pars.UB**
Optional vectors of lower and upper bounds, respectively, used by the "L-BFGS-B" method option in the optim function for the global parameter estimation. Each vector must be of length three, containing values for tausq, sigmasq, and nu. Default for global.pars.LB is rep(1e-05,3); default for global.pars.UB is c(4*resid.var,4*resid.var,100), where resid.var is the residual variance from using lm with mean.model.

**local.ini.pars**
Optional vector of initial values used by the "L-BFGS-B" method option in the optim function for the local parameter estimation. The vector must be of length five, containing values for lam1, lam2, tausq, sigmasq, and nu. Defaults to c(max.distance/10,max.distance/10,0.1*resid.var,0.9*resid.var,1), where max.distance is the maximum interpoint distance of the observed data and resid.var is the residual variance from using lm with mean.model.

**global.ini.pars**
Optional vector of initial values used by the "L-BFGS-B" method option in the optim function for the global parameter estimation. The vector must be of length three, containing values for tausq, sigmasq, and nu. Defaults to c(0.1*resid.var,0.9*resid.var,1), where resid.var is the residual variance from using lm with mean.model.
Value

A "NS convo" object, with the following components:

- `mc.locations`: Mixture component locations used for the simulated data.
- `mc.kernels`: Mixture component kernel matrices used for the simulated data.
- `MLEs.save`: Table of local maximum likelihood estimates for each mixture component location.
- `kernel.ellipses`: N.obs x 2 x 2 array, containing the kernel matrices corresponding to each of the simulated values.
- `data`: Observed data values.
- `beta.GLS`: Generalized least squares estimates of beta, the mean coefficients. For `ns.mean = FALSE`, this is a vector (containing the global mean coefficients); for `ns.mean = TRUE`, this is a matrix (one column for each mixture component location).
- `beta.cov`: Covariance matrix of the generalized least squares estimate of beta. For `ns.mean = FALSE`, this is a matrix (containing the covariance of the global mean coefficients); for `ns.mean = TRUE`, this is an array (one matrix for each mixture component location).
- `Mean.coefs`: "Regression table" for the mean coefficient estimates, listing the estimate, standard error, and t-value (for `ns.mean = FALSE` only).
- `tausq.est`: Estimate of tausq (nugget variance), either scalar (when `ns.nugget = "FALSE"`) or a vector of length N (when `ns.nugget = "TRUE"`), which contains the estimated nugget variance for each observation location.
- `sigmasq.est`: Estimate of sigmasq (process variance), either scalar (when `ns.variance = "FALSE"`) or a vector of length N (when `ns.variance = "TRUE"`), which contains the estimated process variance for each observation location.
- `beta.est`: Estimate of beta (mean coefficients), either a vector (when `ns.mean = "FALSE"`) or a matrix with N rows (when `ns.mean = "TRUE"`), each row of which contains the estimated (smoothed) mean coefficients for each observation location.
- `kappa.MLE`: Scalar maximum likelihood estimate for kappa (when applicable).
- `Cov.mat`: Estimated covariance matrix (N.obs x N.obs) using all relevant parameter estimates.
- `Cov.mat.chol`: Cholesky of `Cov.mat` (i.e., `chol(Cov.mat)`), the estimated covariance matrix (N.obs x N.obs).
- `cov.model`: String; the correlation model used for estimation.
- `ns.nugget`: Logical, indicating if the nugget variance was estimated as spatially-varying (TRUE) or constant (FALSE).
- `ns.variance`: Logical, indicating if the process variance was estimated as spatially-varying (TRUE) or constant (FALSE).
- `fixed.nugg2.var`: N x N matrix with the fixed variance/covariance for the second (measurement error) nugget term (defaults to zero).
- `coords`: N x 2 matrix of observation locations.
global.loglik  Scalar value of the maximized likelihood from the global optimization (if available).

Xmat  Design matrix, obtained from using \texttt{lm} with mean.model.

lambda.w  Tuning parameter for the weight function.

fix.kappa  Logical, indicating if kappa was fixed (TRUE) or estimated (FALSE).

kappa  Scalar; fixed value of kappa.

Examples

```r
## Not run:
# Using white noise data
fit.model <- NSconvo_fit( coords = cbind( runif(100), runif(100)),
data = rnorm(100), fit.radius = 0.4, N.mc = 4 )
## End(Not run)
```

**NSconvo_sim**  
*Simulate data from the nonstationary model.*

**Description**

\texttt{NSconvo_sim} simulates data from the nonstationary model, given mixture component kernel matrices. The function requires either a mixture component kernel object, from the function \texttt{f.mc.kernels()}, or a direct specification of the mixture component locations and mixture component kernels.

**Usage**

```r
NSconvo_sim(
  grid = TRUE,
y.min = 0,
y.max = 5,
x.min = 0,
x.max = 5,
N.obs = 20^2,
sim.locations = NULL,
mc.kernels.obj = NULL,
mc.kernels = NULL,
mc.locations = NULL,
lambda.w = NULL,
tausq = 0.1,
sigmasq = 1,
beta.coefs = 4,
kappa = NULL,
covariates = rep(1, N.obs),
cov.model = "exponential"
)
```
**Arguments**

- **grid** Logical; indicates if the simulated data should fall on a grid (TRUE) or not (FALSE).
- **y.min** Lower bound for the y-coordinate axis.
- **y.max** Upper bound for the y-coordinate axis.
- **x.min** Lower bound for the x-coordinate axis.
- **x.max** Upper bound for the x-coordinate axis.
- **N.obs** Number of simulated data values.
- **sim.locations** Optional N.obs x 2 matrix; allows the user to specify the locations of the simulated data.
- **mc.kernels.obj** Object from the f_mc_kernels function.
- **mc.kernels** Optional specification of mixture component kernel matrices.
- **mc.locations** Optional specification of mixture component locations.
- **lambda.w** Scalar; tuning parameter for the weight function.
- **tausq** Scalar; true nugget variance.
- **sigmasq** Scalar; true process variance.
- **beta.coefs** Vector of true regression coefficients. Length must match the number of columns in covariates.
- **kappa** Scalar; true smoothness.
- **covariates** Matrix with N.obs rows, corresponding to covariate information for each of the simulated values.
- **cov.model** A string specifying the model for the correlation function; defaults to "exponential". Options available in this package are: "exponential", "matern", and "gaussian".

**Value**

A list with the following components:

- **sim.locations** Matrix of locations for the simulated values.
- **mc.locations** Mixture component locations used for the simulated data.
- **mc.kernels** Mixture component kernel matrices used for the simulated data.
- **kernel.ellipses** N.obs x 2 x 2 array, containing the kernel matrices corresponding to each of the simulated values.
- **Cov.mat** True covariance matrix (N.obs x N.obs) corresponding to the simulated data.
- **sim.data** Simulated data values.
- **lambda.w** Tuning parameter for the weight function.
Examples

## Not run:
NSconvo_sim( grid = TRUE, y.min = 0, y.max = 5, x.min = 0, 
x.max = 5, N.obs = 20^2, sim.locations = NULL, mc.kernels.obj = NULL, 
mc.kernels = NULL, mc.locations = NULL, lambda.w = NULL, 
tausq = 0.1, sigmasq = 1, beta.coefs = 4, kappa = NULL, 
covariates = rep(1,N.obs), cov.model = "exponential" )

## End(Not run)

plot.Aniso

Plot of the estimated correlations from the stationary model.

Description

This function plots the estimated correlation between a reference point and all other prediction locations.

Usage

## S3 method for class 'Aniso'
plot(x, ref.loc = NULL, all.pred.locs = NULL, grid = TRUE, ...)

Arguments

x An "Aniso" object, from Aniso_fit().
ref.loc Vector of length 2; the reference location.
all.pred.locs A matrix of all prediction locations.
grid Logical; indicates if the all.pred.locs are on a rectangular grid (TRUE) or not (FALSE).
...
Arguments passed to plot functions.

Value

A plot of either the estimated ellipses or estimated correlation is printed.

Examples

## Not run:
plot.Aniso( Aniso.object )

## End(Not run)
plot.NSconvo

**Plot from the nonstationary model.**

**Description**

This function plots either the estimated anisotropy ellipses for each of the mixture component locations or the estimated correlation between a reference point and all other prediction locations.

**Usage**

```r
## S3 method for class 'NSconvo'
plot(
  x,
  plot.ellipses = TRUE,
  fit.radius = NULL,
  aniso.mat = NULL,
  true.mc = NULL,
  ref.loc = NULL,
  all.pred.locs = NULL,
  grid = TRUE,
  true.col = 1,
  aniso.col = 4,
  ns.col = 2,
  plot.mc.locs = TRUE,
  ...
)
```

**Arguments**

- `x`: A "NSconvo" object, from NSconvo_fit().
- `plot.ellipses`: Logical; indicates whether the estimated ellipses should be plotted (TRUE) or estimated correlations (FALSE).
- `fit.radius`: Scalar; defines the fit radius used for the local likelihood estimation.
- `aniso.mat`: 2 x 2 matrix; contains the estimated anisotropy ellipse from the stationary model (for comparison).
- `true.mc`: The true mixture component ellipses, if known.
- `ref.loc`: Vector of length 2; the reference location.
- `all.pred.locs`: A matrix of all prediction locations.
- `grid`: Logical; indicates if the all.pred.locs are on a rectangular grid (TRUE) or not (FALSE).
- `true.col`: Color value for the true mixture component ellipses (if plotted).
- `aniso.col`: Color value for the anisotropy ellipse (if plotted).
- `ns.col`: Color value for the mixture component ellipses.
- `plot.mc.locs`: Logical; indicates whether the mixture component locations should be plotted (TRUE) or not (FALSE).
- `...`: Other options passed to plot.
Value

A plot of either the estimated ellipses or estimated correlation is printed.

Examples

```r
## Not run:
plot.NSconvo( NSconvo.object )
## End(Not run)
```

predict.Aniso

Obtain predictions at unobserved locations for the stationary spatial model.

Description

predict.Aniso calculates the kriging predictor and corresponding standard errors at unmonitored sites.

Usage

```r
## S3 method for class 'Aniso'
predict(
  object,
  pred.coords,
  pred.covariates = NULL,
  pred.fixed.nugg2.var = NULL,
  ...
)
```

Arguments

- `object` An "Aniso" object, from Aniso_fit.
- `pred.coords` Matrix of locations where predictions are required.
- `pred.covariates` Matrix of covariates for the prediction locations, NOT including an intercept. The number of columns for this matrix must match the design matrix from mean.model in NSconvo_fit. Defaults to an intercept only.
- `pred.fixed.nugg2.var` An optional vector or matrix describing the variance/covariance a fixed second nugget term (corresponds to fixed.nugg2.var in Aniso_fit; often useful if conducting prediction for held-out data). Defaults to zero.
- `...` additional arguments affecting the predictions produced.
predict.NSconvo

Value

A list with the following components:

- pred.means: Vector of the kriging predictor, for each location in `pred.coords`.
- pred.SDs: Vector of the kriging standard errors, for each location in `pred.coords`.

Examples

```r
## Not run:
pred.S <- predict( Aniso.obj, 
pred.coords = cbind(runif(300),runif(300)) )
## End(Not run)
```

---

predict.NSconvo

Obtain predictions at unobserved locations for the nonstationary spatial model.

Description

`predict.NSconvo` calculates the kriging predictor and corresponding standard errors at unmonitored sites.

Usage

```r
## S3 method for class 'NSconvo'
predict( 
  object, 
  pred.coords, 
  pred.covariates = NULL, 
  pred.fixed.nugg2.var = NULL, 
  ... 
)
```

Arguments

- `object`: A "NSconvo" object, from `NSconvo_fit`.
- `pred.coords`: Matrix of locations where predictions are required.
- `pred.covariates`: Matrix of covariates for the prediction locations, NOT including an intercept. The number of columns for this matrix must match the design matrix from `mean.model` in `NSconvo_fit`. Defaults to an intercept only.
- `pred.fixed.nugg2.var`: An optional vector or matrix describing the variance/covariance a fixed second nugget term (corresponds to `fixed.nugg2.var` in `NSconvo_fit`; often useful if conducting prediction for held-out data). Defaults to zero.
- `...`: Additional arguments affecting the predictions produced.
Value

A list with the following components:

pred.means Vector of the kriging predictor, for each location in pred.coords.

pred.SDs Vector of the kriging standard errors, for each location in pred.coords.

Examples

```r
## Not run:
pred.NS <- predict( NSconvo.obj,
pred.coords = matrix(c(1,1), ncol=2),
pred.covariates = matrix(c(1,1), ncol=2) )
## End(Not run)
```

---

**simdata**

*Simulated nonstationary dataset*

Description

A data set containing the necessary components to fit the nonstationary spatial model, simulated from the true model.

Usage

simdata

Format

A list with the following objects:

- **sim.locations** A matrix of longitude/latitude coordinates of the simulated locations.
- **mc.locations** A matrix of longitude/latitude coordinates of the mixture component locations.
- **mc.kernel** A three-dimensional array, containing the true 2 x 2 kernel covariance matrices for each mixture component location.
- **kernel.ellipses** A three-dimensional array, containing the true 2 x 2 kernel covariance matrices for each simulated location.
- **sim.data** A matrix of the simulated data; each of the ten columns correspond to an independent and identically distributed replicate.
- **lambda.w** Scalar; the value of the tuning parameter used in the weight function.
- **holdout.index** Vector; indicates which of the simulated locations should be used in the hold-out sample.
**summary.Aniso**  
*Summarize the stationary model fit.*

**Description**

`summary.Aniso` prints relevant output from the model fitting procedure.

**Usage**

```r
## S3 method for class 'Aniso'
summary(object, ...)  
```

**Arguments**

- `object`  
  An "Aniso" object, from `Aniso_fit`.  
- `...`  
  additional arguments affecting the summary produced.

**Value**

Text containing the model fitting results.

**Examples**

```r
## Not run:
summary.Aniso( Aniso.object )  
## End(Not run)
```

---

**summary.NSconvo**  
*Summarize the nonstationary model fit.*

**Description**

`summary.NSconvo` prints relevant output from the model fitting procedure.

**Usage**

```r
## S3 method for class 'NSconvo'
summary(object, ...)  
```

**Arguments**

- `object`  
  A "NSconvo" object, from `NSconvo_fit`.  
- `...`  
  additional arguments affecting the summary produced.
Value

Text containing the model fitting results.

Examples

```r
## Not run:
summary.NSconvo( NSconvo.object )

## End(Not run)
```

US.mc.grids

*Mixture component grids for the western United States*

Description

A list of two mixture component grids for fitting the nonstationary model to the western United States precipitation data.

Usage

US.mc.grids

Format

A list with two elements:

- **Element 1** Coarse mixture component grid.
- **Element 2** Fine mixture component grid.

US.prediction.locs

*Prediction locations for the western United States*

Description

A matrix with two columns containing a fine grid of locations for which to make a filled-in prediction map for the western United States.

Usage

US.prediction.locs

Format

A matrix with two columns:

- **Column 1** Longitude of the prediction grid.
- **Column 2** Latitude of the prediction grid.
Annual precipitation measurements from the western United States, 1997

Description

A data set containing the annual precipitation for 1270 locations in the western United States.

Usage

USprecip97

Format

A data frame with the following variables:

- longitude Longitude of the monitoring site.
- latitude Latitude of the monitoring site.
- annual.ppt Annual precipitation for the monitoring site, in millimeters.
- log.annual.ppt Annual precipitation for the monitoring site, in log millimeters.

Source

http://www.image.ucar.edu/GSP/Data/US.monthly.met/
Index

* datasets
  simdata, 26
  US.mc.grids, 28
  US.prediction.locs, 28
  USprecip97, 29

Aniso_fit, 2

cov.spatial, 5

evaluate.CV, 5

f_mc_kernels, 6, 21
formula, 3, 17

kernel_cov, 7

lm, 3, 4, 18, 20

make_global_loglik1, 8
make_global_loglik1_kappa, 9
make_global_loglik2, 9
make_global_loglik2_kappa, 10
make_global_loglik3, 11
make_global_loglik3_kappa, 12
make_global_loglik4_kappa, 13
make_local_lik, 14
mc_N, 15

NSconvo_fit, 16, 24, 25
NSconvo_sim, 20

optim, 3, 18

plot.Aniso, 22
plot.NSconvo, 23
predict.Aniso, 24
predict.NSconvo, 25

simdata, 26
summary.Aniso, 27
summary.NSconvo, 27