Package 'CircSpaceTime'

October 12, 2022

```
Type Package
Title Spatial and Spatio-Temporal Bayesian Model for Circular Data
Version 0.9.0
BugReports https://github.com/santoroma/CircSpaceTime
Description Implementation of Bayesian models for spatial and spatio-temporal
      interpolation of circular data using Gaussian Wrapped and Gaussian Projected distributions.
      We developed the methods described in Jona Lasinio G. et al. (2012) <doi:10.1214/12-aoas576>,
      Wang F. et al. (2014) <doi:10.1080/01621459.2014.934454> and
      Mastrantonio G. et al. (2016) <doi:10.1007/s11749-015-0458-y>.
License GPL-3
Encoding UTF-8
LazyData true
URL https://github.com/santoroma/CircSpaceTime
Suggests foreach, iterators, parallel, doParallel, gridExtra
LinkingTo Rcpp, RcppArmadillo, RInside
Imports Rcpp (>= 0.12.14), circular, RInside, coda, ggplot2
RoxygenNote 6.1.0
NeedsCompilation yes
Author Giovanna Jona Lasinio [aut] (<a href="https://orcid.org/0000-0001-8912-5018">https://orcid.org/0000-0001-8912-5018</a>),
      Gianluca Mastrantonio [aut] (<a href="https://orcid.org/0000-0002-2963-6729">https://orcid.org/0000-0002-2963-6729</a>),
      Mario Santoro [aut, cre] (<a href="https://orcid.org/0000-0001-6626-9430">https://orcid.org/0000-0001-6626-9430</a>)
Maintainer Mario Santoro <santoro.ma@gmail.com>
Depends R (>= 2.10)
Repository CRAN
Date/Publication 2019-06-06 15:12:15 UTC
```

2 APEcirc

R topics documented:

	APEcirc																															
	april										•					•			•							•		•			 	5
	CircSpaceTime																														 . (5
	ConvCheck .																														 . (5
	CRPScirc																														 	3
	may																														 10)
	ProjKrigSp .																														 11	1
	ProjKrigSpTi																														 14	4
	ProjSp																														 17	7
	ProjSpTi																														 21	1
	rose_diag																															
	WrapKrigSp .																															
	WrapKrigSpTi																															
	WrapSp																															
	WrapSpTi																															
Index																															38	3
																																_
APEc:	irc		A_1	ver	rag	зe	P	re	di	cti	01	n E	Eri	ror	·fe	or	ci	rc	ul	ar	· V	ar	ia	bl	es.							

Description

APEcirc computes the average prediction error (APE), defined as the average circular distance across pairs

Usage

```
APEcirc(real, sim, bycol = F)
```

Arguments

real	a vector of the values of the process at the test locations
sim	a matrix with nrow = the test locations and ncol = the number of posterior samples from the posterior distributions by WrapKrigSp WrapKrigSpTi, ProjKrigSp, ProjKrigSpTi
bycol	logical. It is TRUE if the columns of sim represent the observations and the rows the posterior samples, the default value is FALSE.

Value

a list of two elements

ApePoints a vector of APE, one element for each test point Ape the overall mean

APEcirc 3

References

G. Jona Lasinio, A. Gelfand, M. Jona-Lasinio, "Spatial analysis of wave direction data using wrapped Gaussian processes", The Annals of Applied Statistics 6 (2013), 1478-1498

See Also

ProjKrigSp and WrapKrigSp for posterior spatial estimations, ProjKrigSpTi and WrapKrigSpTi for posterior spatio-temporal estimations

Other model performance indices: CRPScirc

```
library(CircSpaceTime)
## functions
rmnorm <- function(n = 1, mean = rep(0, d), varcov){
 d <- if (is.matrix(varcov))</pre>
   ncol(varcov)
 else 1
 z <- matrix(rnorm(n * d), n, d) %*% chol(varcov)</pre>
 y \leftarrow t(mean + t(z))
 return(y)
}
## Simulation
set.seed(1)
n <- 20
### simulate coordinates from a unifrom distribution
coords < cbind(runif(n,0,100), runif(n,0,100)) #spatial coordinates
coordsT \leftarrow sort(runif(n,0,100)) #time coordinates (ordered)
Dist <- as.matrix(dist(coords))</pre>
DistT <- as.matrix(dist(coordsT))</pre>
rho
       <- 0.05 #spatial decay
       <- 0.01 #temporal decay
sep_par <- 0.5 #separability parameter</pre>
sigma2 <- 0.3 # variance of the process
alpha <-c(0.5)
#Gneiting covariance
SIGMA \leftarrow sigma2 * (rhoT * DistT^2 + 1)^{-1} * exp(-rho * Dist/(rhoT * DistT^2 + 1)^{-1})
Y \leftarrow rmnorm(1, rep(alpha, times = n), SIGMA) #generate the linear variable
theta <- c()
## wrapping step
for(i in 1:n) {
 theta[i] <- Y[i] %% (2*pi)
### Add plots of the simulated data
```

4 APEcirc

```
rose_diag(theta)
## use this values as references for the definition of initial values and priors
rho_sp.min <- 3/max(Dist)</pre>
rho_sp.max <- rho_sp.min+0.5</pre>
rho_t.min <- 3/max(DistT)</pre>
rho_t.max <- rho_t.min+0.5</pre>
val <- sample(1:n,round(n*0.2)) #validation set</pre>
set.seed(100)
mod <- WrapSpTi(</pre>
         = theta[-val],
 Х
 coords
           = coords[-val,],
           = coordsT[-val],
 times
 start = list("alpha"
                              = c(.79, .74),
                 "rho_sp"
                              = c(.33, .52),
                 "rho_t"
                             = c(.19, .43),
                 "sigma2"
                             = c(.49, .37),
                 "sep_par" = c(.47, .56),
                 "k"
                           = sample(0,length(theta[-val]), replace = TRUE)),
 priors = list("rho_sp"
                                = c(0.01,3/4), ### uniform prior on this interval
                  "rho_t"
                               = c(0.01,3/4), ### uniform prior on this interval
                  "sep_par" = c(1,1), ### beta prior
                  "sigma2"
                              = c(5,5),## inverse gamma prior with mode=5/6
                  "alpha" = c(0,20) ## wrapped gaussian with large variance
 sd_prop = list( "sigma2" = 0.1, "rho_sp" = 0.1, "rho_t" = 0.1, "sep_par"= 0.1),
         = 7000,
 BurninThin = c(burnin = 3000, thin = 10),
 accept_ratio = 0.234,
 adapt_param = c(start = 1, end = 1000, exp = 0.5),
 n_{chains} = 2 ,
 parallel = FALSE,
 n_{cores} = 1
)
check <- ConvCheck(mod, startit = 1 , thin = 1)</pre>
check$Rhat ## convergence has been reached
## when plotting chains remember that alpha is a circular variable
par(mfrow = c(3,2))
coda::traceplot(check$mcmc)
par(mfrow = c(1,1))
########### Prediction on the validation set
Krig <- WrapKrigSpTi(</pre>
 WrapSpTi_out = mod,
 coords_obs = coords[-val,],
 coords_nobs = coords[val,],
 times_obs = coordsT[-val],
 times_nobs = coordsT[val],
 x_{obs} = theta[-val]
)
### checking the prediction
Wrap_Ape <- APEcirc(theta[val], Krig$Prev_out)</pre>
```

april 5

april April waves.

Description

Four days of waves data on the Adriatic sea in the month of April 2010.

Usage

april

Format

Date Date, format: yyyy-mm-dd

hour Factor w/ 24 levels corresponding to the 24h, format: 00:00

Lon, Lat decimal longitude and latitude

Hm0 Significant wave heights in meters

Dm Direction of waves in degrees (North=0)

state Factor w/ 3 levels "calm", "transition", "storm"

Details

Wave directions and heights are obtained as outputs from a deterministic computer model implemented by Istituto Superiore per la Protezione e la Ricerca Ambientale (ISPRA). The computer model starts from a wind forecast model predicting the surface wind over the entire Mediterranean. The hourly evolution of sea wave spectra is obtained by solving energy transport equations using the wind forecast as input. Wave spectra are locally modified using a source function describing the wind energy, the energy redistribution due to nonlinear wave interactions, and energy dissipation due to wave fracture. The model produces estimates every hour on a grid with 10 x 10 km cells (Inghilesi et al. 2016). The ISPRA dataset has forecasts for a total of 4941 grid points over the Italian Mediterranean. Over the Adriatic Sea area, there are 1494 points.

A list containing 4 data frames each of 35856 rows and 7 columns.

Source

R. Inghilesi, A. Orasi & F. Catini (2016) The ISPRA Mediterranean Coastal Wave Forecasting system: evaluation and perspectives Journal of Operational Oceanography Vol. 9, Iss. sup1 http://www.tandfonline.com/doi/full/10.1080/1755876X.2015.1115635

6 ConvCheck

CircSpaceTime	CircSpaceTime: implementation of Bayesian models, for spatial and spatio-temporal interpolation of circular data.
	spano-temporat interpolation of circular data.

Description

The CircSpaceTime package provides two categories of important functions: Sampling Functions and Posterior (Kriging) Estimation Functions.

CircSpaceTime main functions

WrapSp and ProjSp, for sampling from a spatial Normal Wrapped and Projected, respectively. WrapKrigSp and ProjKrigSp, for posterior estimation on spatial Normal Wrapped and Projected, respectively.

WrapSpTi and ProjSpTi, for sampling from a spatio-temporal Normal Wrapped and Projected, respectively. WrapKrigSpTi and ProjKrigSpTi, for posterior estimation on spatio-temporal Normal Wrapped and Projected, respectively.

Description

ConvCheck returns an mcmc.list (mcmc) to be used with the coda package and the Potential scale reduction factors (Rhat) of the model parameters computed using the gelman.diag function in the coda package

Usage

```
ConvCheck(mod, startit = 15000, thin = 10)
```

Arguments

mod is a list with $m \geq 1$ elements, one for each chain generated using Wrap	mod	is a list with m	> 1 eler	nents, one for	each chain	generated using	WrapSp
---	-----	--------------------	----------	----------------	------------	-----------------	--------

ProjSp, WrapSpTi or ProjSpTi

startit is an integer, the iteration at which the chains start (required to build the mcmc.list)

thin is an integer, the thinning applied to chains

Value

a list of two elements

```
mcmc an mcmc.list (mcmc) to be used with the coda package
```

Rhat the Potential scale reduction factors of the model parameters computed using the gelman.diag function in the coda package

ConvCheck 7

See Also

ProjKrigSp and WrapKrigSp for posterior spatial estimations, and ProjKrigSpTi and WrapKrigSpTi for posterior spatio-temporal estimations

```
library(CircSpaceTime)
## auxiliary function
rmnorm<-function(n = 1, mean = rep(0, d), varcov){}
  d <- if (is.matrix(varcov))</pre>
    ncol(varcov)
  else 1
  z <- matrix(rnorm(n * d), n, d) %*% chol(varcov)</pre>
  y \leftarrow t(mean + t(z))
  return(y)
}
####
# Simulation with exponential spatial covariance function
####
set.seed(1)
n <- 20
coords <- cbind(runif(n,0,100), runif(n,0,100))
Dist <- as.matrix(dist(coords))</pre>
rho
        <- 0.05
sigma2 <- 0.3
alpha <- c(0.5)
SIGMA <- sigma2*exp(-rho*Dist)</pre>
Y <- rmnorm(1,rep(alpha,times=n), SIGMA)
theta <- c()
for(i in 1:n) {
  theta[i] <- Y[i]%%(2*pi)
rose_diag(theta)
#validation set
val <- sample(1:n,round(n*0.1))</pre>
set.seed(12345)
mod <- WrapSp(</pre>
  x = theta[-val],
  coords = coords[-val,],
  start = list("alpha" = c(.36, 0.38),
                  "rho"
                           = c(0.041, 0.052),
                 "sigma2" = c(0.24, 0.32),
                          = rep(0,(n - length(val)))),
 priors = list("rho" = c(0.04, 0.08), #few observations require to be more informative
                  "sigma2" = c(2,1),
                  "alpha" = c(0,10)
```

8 CRPScirc

```
),
 sd_prop = list( "sigma2" = 0.1, "rho" = 0.1),
 iter
          = 1000,
 BurninThin
                = c(burnin = 500, thin = 5),
 accept_ratio = 0.234,
 adapt_param = c(start = 40000, end = 45000, exp = 0.5),
 corr_fun = "exponential",
 kappa_matern = .5,
 parallel = FALSE,
 #With doParallel, bigger iter (normally around 1e6) and n_cores>=2 it is a lot faster
 n_{chains} = 2 ,
 n\_cores = 1
check <- ConvCheck(mod)</pre>
check$Rhat ## close to 1 means convergence has been reached
```

CRPScirc

The Continuous Ranked Probability Score for Circular Variables.

Description

CRPScirc function computes the The Continuous Ranked Probability Score for Circular Variables

Usage

```
CRPScirc(obs, sim, bycol = FALSE)
```

Arguments

obs, a vector of the values of the process at the test locations

a matrix with nrow the test locations and ncol the number of posterior samples from the posterior distributions

bycol, logical. It is TRUE if the columns of sim represent the observations and the rows the posterior samples, the default value is FALSE

Value

```
a list of 2 elements

CRPSvec a vector of CRPS, one element for each test point

CRPS the overall mean
```

References

Grimit, Eric P., Tilmann Gneiting, Veronica J. Berrocal, Nicholas Alexander Johnson. "The Continuous Ranked Probability Score for Circular Variables and its Application to Mesoscale Forecast Ensemble Verification", Q.J.R. Meteorol. Soc. 132 (2005), 2925-2942.

CRPScirc 9

See Also

ProjKrigSp and WrapKrigSp for posterior spatial interpolation, and ProjKrigSpTi and WrapKrigSpTi for posterior spatio-temporal interpolation

Other model performance indices: APEcirc

```
#' library(CircSpaceTime)
## auxiliary function
rmnorm<-function(n = 1, mean = rep(0, d), varcov){}
  d <- if (is.matrix(varcov))</pre>
   ncol(varcov)
  else 1
  z <- matrix(rnorm(n * d), n, d) %*% chol(varcov)</pre>
  y \leftarrow t(mean + t(z))
  return(y)
}
####
# Simulation with exponential spatial covariance function
####
set.seed(1)
n <- 20
coords <- cbind(runif(n,0,100), runif(n,0,100))
Dist <- as.matrix(dist(coords))</pre>
rho
       <- 0.05
sigma2 <- 0.3
alpha <- c(0.5)
SIGMA <- sigma2*exp(-rho*Dist)</pre>
Y <- rmnorm(1,rep(alpha,times=n), SIGMA)
theta <- c()
for(i in 1:n) {
  theta[i] <- Y[i]%%(2*pi)
rose_diag(theta)
#validation set
val <- sample(1:n,round(n*0.1))</pre>
set.seed(12345)
mod <- WrapSp(</pre>
 x = theta[-val],
  coords = coords[-val,],
  start = list("alpha" = c(.36, 0.38),
                 "rho"
                          = c(0.041, 0.052),
                 "sigma2" = c(0.24, 0.32),
                          = rep(0,(n - length(val)))),
 priors = list("rho" = c(0.04, 0.08), #few observations require to be more informative
                  "sigma2" = c(2,1),
                  "alpha" = c(0,10)
```

10 may

```
sd_prop = list( "sigma2" = 0.1, "rho" = 0.1),
 iter
         = 1000,
 BurninThin
                = c(burnin = 500, thin = 5),
 accept_ratio = 0.234,
 adapt_param = c(start = 40000, end = 45000, exp = 0.5),
 corr_fun = "exponential",
 kappa_matern = .5,
 parallel = FALSE,
 #With doParallel, bigger iter (normally around 1e6) and n_cores>=2 it is a lot faster
 n_{chains} = 2 ,
 n\_cores = 1
check <- ConvCheck(mod)</pre>
check$Rhat ## close to 1 means convergence has been reached
## graphical check
par(mfrow = c(3,1))
coda::traceplot(check$mcmc)
par(mfrow = c(1,1))
##### We move to the spatial interpolation
Krig <- WrapKrigSp(</pre>
 WrapSp_out = mod,
 coords_obs = coords[-val,],
 coords_nobs = coords[val,],
 x_{obs} = theta[-val]
)
#### check the quality of the prediction using APE and CRPS
ApeCheck <- APEcirc(theta[val],Krig$Prev_out)</pre>
CrpsCheck <- CRPScirc(theta[val],Krig$Prev_out)</pre>
```

may

May waves.

Description

Four time slices of waves data on the Adriatic sea in the month of May 2010.

Usage

may

Format

```
object each element of the list is one hour of data on the entire area
```

Date Date, format: yyyy-mm-dd

hour Factor w/ 24 levels corresponding to the 24h, format: 00:00

Lon, Lat decimal longitude and latitude

ProjKrigSp 11

```
Hm0 Significant wave heights in meters
```

Dm Direction of waves in degrees (North=0)

state Factor w/ 3 levels "calm", "transition", "storm"

Details

Wave directions and heights are obtained as outputs from a deterministic computer model implemented by Istituto Superiore per la Protezione e la Ricerca Ambientale (ISPRA). The computer model starts from a wind forecast model predicting the surface wind over the entire Mediterranean. The hourly evolution of sea wave spectra is obtained by solving energy transport equations using the wind forecast as input. Wave spectra are locally modified using a source function describing the wind energy, the energy redistribution due to nonlinear wave interactions, and energy dissipation due to wave fracture. The model produces estimates every hour on a grid with 10 x 10 km cells (Inghilesi et al. 2016). The ISPRA dataset has forecasts for a total of 4941 grid points over the Italian Mediterranean. Over the Adriatic Sea area, there are 1494 points.

A list containing 4 data frames each of 1494 rows and 7 columns.

Source

R. Inghilesi, A. Orasi & F. Catini (2016) The ISPRA Mediterranean Coastal Wave Forecasting system: evaluation and perspectives Journal of Operational Oceanography Vol. 9, Iss. sup1 http://www.tandfonline.com/doi/full/10.1080/1755876X.2015.1115635

ProjKrigSp

Kriging using projected normal model.

Description

ProjKrigSp function computes the spatial prediction for circular spatial data using samples from the posterior distribution of the spatial projected normal

Usage

```
ProjKrigSp(ProjSp_out, coords_obs, coords_nobs, x_obs)
```

Arguments

ProjSp_out	the function takes the output of ProjSp function
coords_obs	coordinates of observed locations (in UTM)
coords_nobs	coordinates of unobserved locations (in UTM)
x_obs	observed values in $[0,2\pi)$. If they are not in $[0,2\pi)$, the function will transform the data in the right interval

12 ProjKrigSp

Value

a list of 3 elements

M_out the mean of the associated linear process on the prediction locations coords_nobs (rows) over all the posterior samples (columns) returned by ProjSp

V_out the variance of the associated linear process on the prediction locations coords_nobs (rows) over all the posterior samples (columns) returned by ProjSp

Prev_out the posterior predicted values at the unobserved locations.

References

F. Wang, A. E. Gelfand, "Modeling space and space-time directional data using projected Gaussian processes", Journal of the American Statistical Association, 109 (2014), 1565-1580

G. Mastrantonio, G. Jona Lasinio, A. E. Gelfand, "Spatio-temporal circular models with non-separable covariance structure", TEST 25 (2016), 331-350 https://doi.org/10.1007/s11749-015-0458-y

See Also

ProjSp for spatial sampling from Projected Normal , WrapSp for spatial sampling from Wrapped Normal and WrapKrigSp for spatial interpolation under the wrapped model

Other spatial interpolations: WrapKrigSp

```
library(CircSpaceTime)
## auxiliary function
rmnorm <- function(n = 1, mean = rep(0, d), varcov){}
 d <- if (is.matrix(varcov))</pre>
   ncol(varcov)
 else 1
 z <- matrix(rnorm(n * d), n, d) %*% chol(varcov)</pre>
 y <- t(mean + t(z))
 return(y)
}
####
# Simulation using exponential spatial covariance function
set.seed(1)
n <- 20
coords <- cbind(runif(n,0,100), runif(n,0,100))
Dist <- as.matrix(dist(coords))</pre>
rho
        <- 0.05
        <- 0.2
tau
sigma2 <- 1
alpha <-c(0.5,0.5)
SIGMA <- sigma2*exp(-rho*Dist)
```

ProjKrigSp 13

```
Y <- rmnorm(1,rep(alpha,times=n),
       kronecker(SIGMA, matrix(c( sigma2, sqrt(sigma2)*tau, sqrt(sigma2)*tau, 1 ) , nrow=2 )))
theta <- c()
for(i in 1:n) {
theta[i] \leftarrow atan2(Y[(i-1)*2+2],Y[(i-1)*2+1])
theta <- theta %% (2*pi) #to be sure to have values in (0,2pi)
hist(theta)
rose_diag(theta)
val <- sample(1:n,round(n*0.1))</pre>
#############some useful quantities
rho.min <- 3/max(Dist)</pre>
rho.max <- rho.min+0.5</pre>
set.seed(100)
mod <- ProjSp(</pre>
       = theta[-val],
Х
 coords = coords[-val,],
 start = list("alpha"
                            = c(0.92, 0.18, 0.56, -0.35),
                "rho"
                          = c(0.51, 0.15),
                "tau"
                          = c(0.46, 0.66),
                "sigma2"
                           = c(0.27, 0.3),
                "r"
                         = abs(rnorm( length(theta)) )),
 priors = list("rho"
                            = c(rho.min,rho.max),
                 "tau"
                           = c(-1,1),
                 "sigma2"
                            = c(10,3),
                 "alpha_mu" = c(0, 0),
                 "alpha_sigma" = diag(10,2)
 sd_prop = list("sigma2" = 0.1, "tau" = 0.1, "rho" = 0.1,
                  "sdr" = sample(.05,length(theta), replace = TRUE)),
 iter = 10000,
 BurninThin = c(burnin = 7000, thin = 10),
 accept_ratio = 0.234,
 adapt_param = c(start = 130000, end = 120000, exp = 0.5), #no adaptation
 corr_fun = "exponential",
 kappa_matern = .5,
 n_{chains} = 2 ,
 parallel = TRUE ,
n\_cores = 2
)
# If you don't want to install/use DoParallel
# please set parallel = FALSE. Keep in mind that it can be substantially slower
# How much it takes?
check <- ConvCheck(mod)</pre>
check$Rhat #close to 1 we have convergence
#### graphical check
```

14 ProjKrigSpTi

```
par(mfrow=c(3,2))
coda::traceplot(check$mcmc)

par(mfrow=c(1,1))

# move to prediction once convergence is achieved

Krig <- ProjKrigSp(
    ProjSp_out = mod,
    coords_obs = coords[-val,],
    coords_nobs = coords[val,],
    x_obs = theta[-val]
)

# The quality of prediction can be checked using APEcirc and CRPScirc ape <- APEcirc(theta[val],Krig$Prev_out)
crps <- CRPScirc(theta[val],Krig$Prev_out)</pre>
```

ProjKrigSpTi

#' Spatio temporal interpolation using projected spatial temporal normal model.

Description

ProjKrigSpTi function computes the spatio-temporal prediction for circular space-time data using samples from the posterior distribution of the space-time projected normal model.

Usage

```
ProjKrigSpTi(ProjSpTi_out, coords_obs, coords_nobs, times_obs, times_nobs, x_obs)
```

Arguments

ProjSpTi_out the functions takes the output of ProjSpTi function coords_obs coordinates of observed locations (in UTM) coords_nobs coordinates of unobserved locations (in UTM) numeric vector of observed time coordinates numeric vector of unobserved time coordinates v_obs observed values in $[0, 2\pi)$ If they are not in $[0, 2\pi)$, the function will tranform the data in the right interval

Value

a list of 3 elements

M_out the mean of the associated linear process on the prediction locations coords_nobs (rows) over all the posterior samples (columns) returned by ProjSpTi

V_out the variance of the associated linear process on the prediction locations coords_nobs (rows) over all the posterior samples (columns) returned by ProjSpTi

Prev_out are the posterior predicted values at the unobserved locations.

ProjKrigSpTi 15

References

G. Mastrantonio, G.Jona Lasinio, A. E. Gelfand, "Spatio-temporal circular models with non-separable covariance structure", TEST 25 (2016), 331–350.

F. Wang, A. E. Gelfand, "Modeling space and space-time directional data using projected Gaussian processes", Journal of the American Statistical Association, 109 (2014), 1565-1580

T. Gneiting, "Nonseparable, Stationary Covariance Functions for Space-Time Data", JASA 97 (2002), 590-600

See Also

ProjSpTi to sample the posterior distribution of the spatio-temporal Projected Normal model, WrapSpTi to sample the posterior distribution of the spatio-temporal Wrapped Normal model and WrapKrigSpTi for spatio-temporal interpolation under the same model

```
library(CircSpaceTime)
#### simulated example
## auxiliary functions
rmnorm \leftarrow function(n = 1, mean = rep(0, d), varcov) {
  d <- if (is.matrix(varcov)) {</pre>
    ncol(varcov)
  } else {
    1
  }
  z <- matrix(rnorm(n * d), n, d) %*% chol(varcov)</pre>
  v \leftarrow t(mean + t(z))
  return(y)
}
####
# Simulation using a gneiting covariance function
set.seed(1)
n <- 20
coords <- cbind(runif(n, 0, 100), runif(n, 0, 100))
coordsT <- cbind(runif(n, 0, 100))</pre>
Dist <- as.matrix(dist(coords))</pre>
DistT <- as.matrix(dist(coordsT))</pre>
rho <- 0.05
rhoT <- 0.01
sep_par <- 0.1
sigma2 <- 1
SIGMA \leftarrow sigma2 * (rhoT * DistT^2 + 1)^(-1) * exp(-rho * Dist / (rhoT * DistT^2 + 1)^(sep_par / 2))
tau <- 0.2
Y <- rmnorm(
  1, rep(alpha, times = n),
 kronecker(SIGMA, matrix(c(sigma2, sqrt(sigma2) * tau, sqrt(sigma2) * tau, 1), nrow = 2))
```

16 ProjKrigSpTi

```
theta <- c()
for (i in 1:n) {
 theta[i] <- atan2(Y[(i - 1) * 2 + 2], Y[(i - 1) * 2 + 1])
theta <- theta %% (2 * pi) ## to be sure we have values in (0,2pi)
rose_diag(theta)
############ some useful quantities
rho_sp.min <- 3 / max(Dist)</pre>
rho_sp.max <- rho_sp.min + 0.5</pre>
rho_t.min <- 3 / max(DistT)</pre>
rho_t.max <- rho_t.min + 0.5</pre>
### validation set 20% of the data
val <- sample(1:n, round(n \star 0.2))
set.seed(200)
mod <- ProjSpTi(</pre>
 x = theta[-val],
 coords = coords[-val, ],
 times = coordsT[-val],
 start = list(
    "alpha" = c(0.66, 0.38, 0.27, 0.13),
    "rho_sp" = c(0.29, 0.33),
    "rho_t" = c(0.25, 0.13),
    "sep_par" = c(0.56, 0.31),
    "tau" = c(0.71, 0.65),
    "sigma2" = c(0.47, 0.53),
    "r" = abs(rnorm(length(theta[-val])))
 ),
 priors = list(
    "rho_sp" = c(rho_sp.min, rho_sp.max), # Uniform prior in this interval
    "rho_t" = c(rho_t.min, rho_t.max), # Uniform prior in this interval
    "sep_par" = c(1, 1), # Beta distribution
    "tau" = c(-1, 1), ## Uniform prior in this interval
    "sigma2" = c(10, 3), # inverse gamma
    "alpha_mu" = c(0, 0), ## a vector of 2 elements,
    ## the means of the bivariate Gaussian distribution
    "alpha_sigma" = diag(10, 2) # a 2x2 matrix, the covariance matrix of the
    # bivariate Gaussian distribution,
 ),
 sd_prop = list(
    "sep_par" = 0.1, "sigma2" = 0.1, "tau" = 0.1, "rho_sp" = 0.1, "rho_t" = 0.1,
    "sdr" = sample(.05, length(theta), replace = TRUE)
 ),
 iter = 4000,
 BurninThin = c(burnin = 2000, thin = 2),
 accept_ratio = 0.234,
 adapt_param = c(start = 155000, end = 156000, exp = 0.5),
 n_{chains} = 2,
 parallel = TRUE,
check <- ConvCheck(mod)</pre>
```

```
check$Rhat ### convergence has been reached when the values are close to 1
#### graphical checking
#### recall that it is made of as many lists as the number of chains and it has elements named
#### as the model's parameters
par(mfrow = c(3, 3))
coda::traceplot(check$mcmc)
par(mfrow = c(1, 1))
# once convergence is reached we run the interpolation on the validation set
Krig <- ProjKrigSpTi(</pre>
  ProjSpTi_out = mod,
  coords_obs = coords[-val, ],
  coords_nobs = coords[val, ],
  times_obs = coordsT[-val],
  times_nobs = coordsT[val],
  x_{obs} = theta[-val]
#### checking the prediction
Proj_ape <- APEcirc(theta[val], Krig$Prev_out)</pre>
Proj_crps <- CRPScirc(theta[val],Krig$Prev_out)</pre>
```

ProjSp

Samples from the Projected Normal spatial model

Description

ProjSp produces samples from the posterior distribtion of the spatial projected normal model.

Usage

```
ProjSp(x = x, coords = coords, start = list(alpha = c(1, 1, 0.5,
    0.5), tau = c(0.1, 0.5), rho = c(0.1, 0.5), sigma2 = c(0.1, 0.5), r =
    rep(1, times = length(x))), priors = list(tau = c(8, 14), rho = c(8,
    14), sigma2 = c(), alpha_mu = c(1, 1), alpha_sigma = c()),
    sd_prop = list(sigma2 = 0.5, tau = 0.5, rho = 0.5, sdr = sample(0.05,
    length(x), replace = TRUE)), iter = 1000, BurninThin = c(burnin = 20,
    thin = 10), accept_ratio = 0.234, adapt_param = c(start = 1, end =
    1e+07, exp = 0.9, sdr_update_iter = 50), corr_fun = "exponential",
    kappa_matern = 0.5, n_chains = 2, parallel = FALSE, n_cores = 1)
```

Arguments

X	a vector of n circular data in $[0, 2\pi)$. If they are not in $[0, 2\pi)$, the function will
	transform the data in the right interval
coords	an nx2 matrix with the sites coordinates
start	a list of 4 elements giving initial values for the model parameters. Each elements is a vector with n_chains elements
	is a vector with n_chains elements

• alpha the 2-d vector of the means of the Gaussian bi-variate distribution,

- tau the correlation of the two components of the linear representation,
- rho the spatial decay parameter,
- sigma2 the process variance,
- r the vector of length(x), latent variable

priors a list of 4 elements to define priors for the model parameters:

alpha_mu a vector of 2 elements, the means of the bivariate Gaussian distribution,

alpha_sigma a 2x2 matrix, the covariance matrix of the bivariate Gaussian distribution,

rho vector of 2 elements defining the minimum and maximum of a uniform distribution,

tau vector of 2 elements defining the minimum and maximum of a uniform distribution, with the constraints min(tau) >= -1 and max(tau) <= 1

sigma2 a vector of 2 elements defining the shape and rate of an inverse-gamma distribution.

sd_prop list of 4 elements. To run the MCMC for the rho, tau and sigma2 parameters and

r vector we use an adaptive metropolis and in sd_prop we build a list of initial

guesses for these three parameters and the r vector

iter number of iterations

BurninThin a vector of 2 elements with the burnin and the chain thinning accept_ratio it is the desired acceptance ratio in the adaptive metropolis

adapt_param a vector of 4 elements giving the iteration number at which the adaptation must

start and end. The third element (exp) must be a number in (0,1) is a parameter ruling the speed of changes in the adaptation algorithm, it is recommended to set it close to 1, if it is too small non positive definite matrices may be generated and the program crashes. The last element (sdr_update_iter) must be a positive integer defining every how many iterations there is the update of the sd (vector)

of (vector) r.

corr_fun characters, the name of the correlation function; currently implemented func-

tions are c("exponential", "matern", "gaussian")

kappa_matern numeric, the smoothness parameter of the Matern correlation function, default

is kappa_matern = 0.5 (the exponential function)

n_chains integer, the number of chains to be launched (default is 1, but we recommend to

use at least 2 for model diagnostic)

parallel logical, if the multiple chains must be lunched in parallel (you should install

doParallel package). Default is FALSE

n_cores integer, required if parallel=TRUE, the number of cores to be used in the imple-

mentation. Default value is 1.

Value

it returns a list of n_chains lists each with elements

```
rho,tau, sigma2 vectors with the thinned chains alpha a matrix with nrow=2 and ncol= the length of thinned chains, r a matrix with nrow=length(x) and ncol= the length of thinned chains corr_fun characters with the type of spatial correlation chosen distribution characters, always "ProjSp"
```

References

- G. Mastrantonio , G. Jona Lasinio, A. E. Gelfand, "Spatio-temporal circular models with non-separable covariance structure", TEST 25 (2016), 331–350.
- F. Wang, A. E. Gelfand, "Modeling space and space-time directional data using projected Gaussian processes", Journal of the American Statistical Association, 109 (2014), 1565-1580

See Also

ProjKrigSp for spatial interpolation under the projected normal model, WrapSp for spatial sampling from Wrapped Normal and WrapKrigSp for Kriging estimation

```
library(CircSpaceTime)
## auxiliary function
rmnorm <- function(n = 1, mean = rep(0, d), varcov){
 d <- if (is.matrix(varcov))</pre>
   ncol(varcov)
 else 1
 z <- matrix(rnorm(n * d), n, d) %*% chol(varcov)</pre>
 y \leftarrow t(mean + t(z))
 return(y)
# Simulation using exponential spatial covariance function
####
set.seed(1)
n <- 20
coords <- cbind(runif(n,0,100), runif(n,0,100))
Dist <- as.matrix(dist(coords))</pre>
rho
        <- 0.05
        <- 0.2
tau
sigma2 <- 1
alpha <-c(0.5,0.5)
SIGMA
       <- sigma2*exp(-rho*Dist)
Y <- rmnorm(1,rep(alpha,times=n),
       kronecker(SIGMA, matrix(c( sigma2, sqrt(sigma2)*tau, sqrt(sigma2)*tau, 1 ) , nrow=2 )))
theta <- c()
for(i in 1:n) {
```

```
theta[i] <- atan2(Y[(i-1)*2+2],Y[(i-1)*2+1])
theta <- theta %% (2*pi) #to be sure to have values in (0,2pi)
hist(theta)
rose_diag(theta)
val <- sample(1:n,round(n*0.1))</pre>
#############some useful quantities
rho.min <- 3/max(Dist)</pre>
rho.max <- rho.min+0.5</pre>
set.seed(100)
mod <- ProjSp(</pre>
X
       = theta[-val],
 coords = coords[-val,],
 start = list("alpha"
                           = c(0.92, 0.18, 0.56, -0.35),
                "rho"
                         = c(0.51, 0.15),
                "tau"
                         = c(0.46, 0.66),
                "sigma2"
                          = c(0.27, 0.3),
                         = abs(rnorm( length(theta)) )),
 priors = list("rho"
                           = c(rho.min,rho.max),
                 "tau"
                            = c(-1,1),
                 "sigma2"
                           = c(10,3),
                 "alpha_mu" = c(0, 0),
                 "alpha_sigma" = diag(10,2)
 ) ,
 sd_prop = list("sigma2" = 0.1, "tau" = 0.1, "rho" = 0.1,
                  "sdr" = sample(.05,length(theta), replace = TRUE)),
 iter = 10000,
 BurninThin
              = c(burnin = 7000, thin = 10),
 accept_ratio = 0.234,
 adapt_param = c(start = 130000, end = 120000, exp = 0.5), #no adaptation
 corr_fun = "exponential",
 kappa_matern = .5,
 n_{chains} = 2 ,
 parallel = TRUE ,
 n\_cores = 2
# If you don't want to install/use DoParallel
# please set parallel = FALSE. Keep in mind that it can be substantially slower
# How much it takes?
check <- ConvCheck(mod)</pre>
check$Rhat #close to 1 we have convergence
#### graphical check
par(mfrow=c(3,2))
coda::traceplot(check$mcmc)
par(mfrow=c(1,1))
```

once convergence is achieved move to prediction using ProjKrigSp

ProjSpTi

Samples from the posterior distribution of the Projected Normal spatial model

Description

ProjSpTi produces samples from the posterior distribution of the spatial projected normal model.

Usage

```
ProjSpTi(x = x, coords = coords, times = c(), start = list(alpha =
    c(1, 1, 0.5, 0.5), tau = c(0.1, 0.5), rho_sp = c(0.1, 0.5), rho_t =
    c(0.1, 0.5), sep_par = c(0.1, 0.5), sigma2 = c(0.1, 0.5), r = sample(1,
    length(x), replace = T)), priors = list(tau = c(8, 14), rho_sp = c(8,
    14), rho_t = c(8, 14), sep_par = c(8, 14), sigma2 = c(), alpha_mu = c(1,
    1), alpha_sigma = c()), sd_prop = list(sigma2 = 0.5, tau = 0.5, rho_sp
    = 0.5, rho_t = 0.5, sep_par = 0.5, sdr = sample(0.05, length(x), replace
    = T)), iter = 1000, BurninThin = c(burnin = 20, thin = 10),
    accept_ratio = 0.234, adapt_param = c(start = 1, end = 1e+07, exp =
    0.9, sdr_update_iter = 50), n_chains = 2, parallel = FALSE,
    n_cores = 1)
```

Arguments

Χ

a vector of n circular data in $[0,2\pi)$ If they are not in $[0,2\pi)$, the function will tranform the data in the right interval

coords

an nx2 matrix with the sites coordinates

times

an n vector with the times of

start

a list of 4 elements giving initial values for the model parameters. Each elements is a vector with n_chains elements

- alpha the 2-d vector of the means of the Gaussian bi-variate distribution,
- tau the correlation of the two components of the linear representation,
- rho_sp the spatial decay parameter,
- rho_t the temporal decay parameter,
- sigma2 the process variance,
- sep_par the separation parameter,
- r the vector of length(x), latent variable

priors

a list of 7 elements to define priors for the model parameters:

alpha_mu a vector of 2 elements, the means of the bivariate Gaussian distribution

alpha_sigma a 2x2 matrix, the covariance matrix of the bivariate Gaussian distribution,

rho_sp a vector of 2 elements defining the minimum and maximum of a uniform distribution,

rho_t a vector of 2 elements defining the minimum and maximum of a uniform distribution,

tau vector of 2 elements defining the minimum and maximum of a uniform distribution with the constraints min(tau) >= -1 and max(tau) <= 1,

sep_par a vector of 2 elements defining the two parameters of a beta distribution,

sigma2 a vector of 2 elements defining the shape and rate of an inverse-gamma distribution,

sd_prop = list of 4 elements. To run the MCMC for the rho_sp, tau and sigma2 parameters

and r vector we use an adaptive metropolis and in sd_prop we build a list of

initial guesses for these three parameters and the r vector

iter iter number of iterations

BurninThin a vector of 2 elements with the burnin and the chain thinning accept_ratio it is the desired acceptance ratio in the adaptive metropolis

adapt_param a vector of 4 elements giving the iteration number at which the adaptation must

start and end. The third element (exp) must be a number in (0,1) is a parameter ruling the speed of changes in the adaptation algorithm, it is recommended to set it close to 1, if it is too small non positive definite matrices may be generated and the program crashes. The last element (sdr_update_iter) must be a positive integer defining every how many iterations there is the update of the sd (vector)

of (vector) r.

n_chains integer, the number of chains to be launched (default is 1, but we recommend to

use at least 2 for model diagnostic)

parallel logical, if the multiple chains must be lunched in parallel (you should install

doParallel package). Default is FALSE

n_cores integer, required if parallel=TRUE, the number of cores to be used in the imple-

mentation. Default value is 1.

Value

it returns a list of n_chains lists each with elements

tau, rho_sp, rho_t, sigma2 vectors with the thinned chains

alpha a matrix with nrow=2 and ncol= the length of thinned chains

r a matrix with nrow=length(x) and ncol= the length of thinned chains

References

G. Mastrantonio, G.Jona Lasinio, A. E. Gelfand, "Spatio-temporal circular models with non-separable covariance structure", TEST 25 (2016), 331–350.

F. Wang, A. E. Gelfand, "Modeling space and space-time directional data using projected Gaussian processes", Journal of the American Statistical Association, 109 (2014), 1565-1580

T. Gneiting, "Nonseparable, Stationary Covariance Functions for Space-Time Data", JASA 97 (2002), 590-600

See Also

ProjKrigSpTi for spatio-temporal prediction under the spatio-temporal projected normal model, WrapSpTi to sample from the posterior distribution of the spatio-temporal Wrapped Normal model and WrapKrigSpTi for spatio-temporal prediction under the same model

Other spatio-temporal models: WrapSpTi

```
library(CircSpaceTime)
#### simulated example
## auxiliary functions
rmnorm <- function(n = 1, mean = rep(0, d), varcov) {
  d <- if (is.matrix(varcov)) {</pre>
    ncol(varcov)
  } else {
    1
  }
  z <- matrix(rnorm(n * d), n, d) %*% chol(varcov)</pre>
  y \leftarrow t(mean + t(z))
  return(y)
}
####
# Simulation using a gneiting covariance function
####
set.seed(1)
n <- 20
coords <- cbind(runif(n, 0, 100), runif(n, 0, 100))
coordsT <- cbind(runif(n, 0, 100))</pre>
Dist <- as.matrix(dist(coords))</pre>
DistT <- as.matrix(dist(coordsT))</pre>
rho <- 0.05
rhoT <- 0.01
sep_par <- 0.1
sigma2 <- 1
alpha \leftarrow c(0.5)
SIGMA \leftarrow sigma2 * (rhoT * DistT^2 + 1)^{-1} * exp(-rho * Dist / (rhoT * DistT^2 + 1)^(sep_par / 2))
tau <- 0.2
Y <- rmnorm(
  1, rep(alpha, times = n),
 kronecker(SIGMA, matrix(c(sigma2, sqrt(sigma2) * tau, sqrt(sigma2) * tau, 1), nrow = 2))
theta <- c()
for (i in 1:n) {
  theta[i] <- atan2(Y[(i - 1) * 2 + 2], Y[(i - 1) * 2 + 1])
theta <- theta %% (2 * pi) ## to be sure we have values in (0,2pi)
rose_diag(theta)
########### some useful quantities
```

```
rho_sp.min <- 3 / max(Dist)</pre>
rho_sp.max <- rho_sp.min + 0.5</pre>
rho_t.min <- 3 / max(DistT)</pre>
rho_t.max <- rho_t.min + 0.5
### validation set 20% of the data
val <- sample(1:n, round(n * 0.2))
set.seed(200)
mod <- ProjSpTi(</pre>
 x = theta[-val],
 coords = coords[-val, ],
 times = coordsT[-val],
 start = list(
    "alpha" = c(0.66, 0.38, 0.27, 0.13),
    "rho_sp" = c(0.29, 0.33),
    "rho_t" = c(0.25, 0.13),
    "sep_par" = c(0.56, 0.31),
    "tau" = c(0.71, 0.65),
    "sigma2" = c(0.47, 0.53),
    "r" = abs(rnorm(length(theta[-val])))
 ),
 priors = list(
    "rho_sp" = c(rho_sp.min, rho_sp.max), # Uniform prior in this interval
    "rho_t" = c(rho_t.min, rho_t.max), \# Uniform prior in this interval
    "sep_par" = c(1, 1), # Beta distribution
    "tau" = c(-1, 1), ## Uniform prior in this interval
    "sigma2" = c(10, 3), # inverse gamma
    "alpha_mu" = c(0, 0), ## a vector of 2 elements,
    ## the means of the bivariate Gaussian distribution
    "alpha_sigma" = diag(10, 2) # a 2x2 matrix, the covariance matrix of the
    # bivariate Gaussian distribution,
 sd_prop = list(
    "sep_par" = 0.1, "sigma2" = 0.1, "tau" = 0.1, "rho_sp" = 0.1, "rho_t" = 0.1,
    "sdr" = sample(.05, length(theta), replace = TRUE)
 ),
 iter = 4000,
 BurninThin = c(burnin = 2000, thin = 2),
 accept_ratio = 0.234,
 adapt_param = c(start = 155000, end = 156000, exp = 0.5),
 n_{chains} = 2,
 parallel = TRUE,
)
check <- ConvCheck(mod)</pre>
check$Rhat ### convergence has been reached when the values are close to 1
#### graphical checking
#### recall that it is made of as many lists as the number of chains and it has elements named
#### as the model's parameters
par(mfrow = c(3, 3))
coda::traceplot(check$mcmc)
par(mfrow = c(1, 1))
# move to prediction once convergence is achieved using ProjKrigSpTi
```

rose_diag 25

rose	dia	0

Rose diagram in ggplot2 inspired from rose.diag in package circular.

Description

Rose diagram in ggplot2 inspired from rose.diag in package circular.

Usage

```
rose_diag(x, bins = 15, color = "red", alpha = 1, start = 0,
  add = FALSE, template = "rad", direction = NULL)
```

Arguments

x	a vector of circular coordinates in radiants $[0, 2\pi)$.
bins	number of bins
color	color of the line and of the fill
alpha	transparency
start	the starting angle of the 0 (the North)
add	add the rose_diag to an existing ggplot2 plot
template	radiants or wind rose. the values are c("rad", "wind_rose") default is "rad"
direction	1, clockwise; -1, anticlockwise. For template = "rad" direction is -1 while for template = "wind rose" direction is 1.

Value

The plot in ggplot2 format.

```
library(CircSpaceTime)
x <- circular::rwrappedstable(200, index = 1.5, skewness = .5)
x1 <- circular::rwrappedstable(200, index = 2, skewness = .5)
x2 <- circular::rwrappedstable(200, index = 0.5, skewness = 1)
rose_diag(x, bins = 15, color = "green")
rose_diag(x1, bins = 15, color = "blue", alpha = .5, add = TRUE)
rose_diag(x2, bins = 15, color = "red", alpha = .5, add = TRUE)</pre>
```

26 WrapKrigSp

WrapKrigSp	Spatial interpolation using wrapped normal model.

Description

WrapKrigSp function computes the spatial prediction for circular spatial data using samples from the posterior distribution of the spatial wrapped normal

Usage

```
WrapKrigSp(WrapSp_out, coords_obs, coords_nobs, x_obs)
```

Arguments

WrapSp_out the functions takes the output of WrapSp function coords_obs coordinates of observed locations (in UTM) coords_nobs coordinates of unobserved locations (in UTM) observed values

Value

a list of 3 elements

M_out the mean of the associated linear process on the prediction locations coords_nobs (rows) over all the posterior samples (columns) returned by WrapSp

V_out the variance of the associated linear process on the prediction locations coords_nobs (rows) over all the posterior samples (columns) returned by WrapSp

Prev_out the posterior predicted values at the unobserved locations.

Implementation Tips

To facilitate the estimations, the observations x are centered around pi, and the posterior samples of x and posterior mean are changed back to the original scale

References

G. Jona-Lasinio, A. E. Gelfand, M. Jona-Lasinio, "Spatial analysis of wave direction data using wrapped Gaussian processes", The Annals of Applied Statistics, 6 (2012), 1478-1498

See Also

WrapSp for spatial sampling from Wrapped Normal , ProjSp for spatial sampling from Projected Normal and ProjKrigSp for Kriging estimation

Other spatial interpolations: ProjKrigSp

WrapKrigSp 27

```
library(CircSpaceTime)
## auxiliary function
rmnorm<-function(n = 1, mean = rep(0, d), varcov){}
  d <- if (is.matrix(varcov))</pre>
   ncol(varcov)
  else 1
  z <- matrix(rnorm(n * d), n, d) %*% chol(varcov)</pre>
  y \leftarrow t(mean + t(z))
  return(y)
}
####
# Simulation with exponential spatial covariance function
####
set.seed(1)
n <- 20
coords <- cbind(runif(n,0,100), runif(n,0,100))
Dist <- as.matrix(dist(coords))</pre>
rho
        <- 0.05
sigma2 <- 0.3
alpha <-c(0.5)
SIGMA <- sigma2*exp(-rho*Dist)</pre>
Y <- rmnorm(1,rep(alpha,times=n), SIGMA)
theta <- c()
for(i in 1:n) {
  theta[i] <- Y[i]%%(2*pi)
}
rose_diag(theta)
#validation set
val <- sample(1:n,round(n*0.1))</pre>
set.seed(12345)
mod <- WrapSp(</pre>
          = theta[-val],
 coords = coords[-val,],
  start = list("alpha"
                             = c(.36, 0.38),
                           = c(0.041, 0.052),
                 "rho"
                 "sigma2" = c(0.24, 0.32),
                          = rep(0,(n - length(val)))),
 priors = list("rho"
                        = c(0.04, 0.08), #few observations require to be more informative
                  "sigma2"
                             = c(2,1),
                  "alpha" = c(0,10)
  ),
  sd_prop = list( "sigma2" = 0.1, "rho" = 0.1),
  iter
         = 1000,
  BurninThin = c(burnin = 500, thin = 5),
  accept_ratio = 0.234,
  adapt_param = c(start = 40000, end = 45000, exp = 0.5),
```

28 WrapKrigSpTi

```
corr_fun = "exponential",
 kappa_matern = .5,
 parallel = FALSE,
 #With doParallel, bigger iter (normally around 1e6) and n_cores>=2 it is a lot faster
 n_{chains} = 2 ,
 n\_cores = 1
)
check <- ConvCheck(mod)</pre>
check$Rhat ## close to 1 means convergence has been reached
## graphical check
par(mfrow = c(3,1))
coda::traceplot(check$mcmc)
par(mfrow = c(1,1))
##### We move to the spatial interpolation
Krig <- WrapKrigSp(</pre>
 WrapSp_out = mod,
 coords_obs = coords[-val,],
 coords_nobs = coords[val,],
 x_{obs} = theta[-val]
)
\#\#\# check the quality of the prediction using APE and CRPS
ApeCheck <- APEcirc(theta[val],Krig$Prev_out)</pre>
CrpsCheck <- CRPScirc(theta[val],Krig$Prev_out)</pre>
```

WrapKrigSpTi

Prediction using wrapped normal spatio-temporal model.

Description

WrapKrigSpTi function computes the spatio-temporal prediction for circular space-time data using samples from the posterior distribution of the space-time wrapped normal model

Usage

```
WrapKrigSpTi(WrapSpTi_out, coords_obs, coords_nobs, times_obs, times_nobs,
  x_obs)
```

Arguments

```
WrapSpTi_out the functions takes the output of WrapSpTi function coords_obs coordinates of observed locations (in UTM)

coords_nobs coordinates of unobserved locations (in UTM)

times_obs numeric vector of observed time coordinates

times_nobs observed values
```

WrapKrigSpTi 29

Value

```
a list of 3 elements
```

M_out the mean of the associated linear process on the prediction locations coords_nobs (rows) over all the posterior samples (columns) returned by WrapSpTi

V_out the variance of the associated linear process on the prediction locations coords_nobs (rows) over all the posterior samples (columns) returned by WrapSpTi

Prev_out the posterior predicted values at the unobserved locations

Implementation Tips

To facilitate the estimations, the observations x are centered around π . Posterior samples of x at the predictive locations and posterior mean are changed back to the original scale

References

- G. Mastrantonio, G. Jona Lasinio, A. E. Gelfand, "Spatio-temporal circular models with non-separable covariance structure", TEST 25 (2016), 331–350
- T. Gneiting, "Nonseparable, Stationary Covariance Functions for Space-Time Data", JASA 97 (2002), 590-600

See Also

WrapSpTi spatio-temporal sampling from Wrapped Normal, ProjSpTi for spatio-temporal sampling from Projected Normal and ProjKrigSpTi for Kriging estimation

```
library(CircSpaceTime)
## functions
rmnorm <- function(n = 1, mean = rep(0, d), varcov){
 d <- if (is.matrix(varcov))</pre>
   ncol(varcov)
 z <- matrix(rnorm(n * d), n, d) %*% chol(varcov)</pre>
 y \leftarrow t(mean + t(z))
 return(y)
## Simulation
set.seed(1)
n <- 20
### simulate coordinates from a unifrom distribution
coords <- cbind(runif(n,0,100), runif(n,0,100)) \#spatial coordinates
coordsT <- sort(runif(n,0,100)) #time coordinates (ordered)</pre>
Dist <- as.matrix(dist(coords))</pre>
DistT <- as.matrix(dist(coordsT))</pre>
```

30 WrapKrigSpTi

```
<- 0.05 #spatial decay
rho
rhoT
        <- 0.01 #temporal decay
sep_par <- 0.5 #separability parameter</pre>
sigma2 <- 0.3 # variance of the process
alpha <-c(0.5)
#Gneiting covariance
SIGMA \leftarrow sigma2 * (rhoT * DistT^2 + 1)^{-1} * exp(-rho * Dist/(rhoT * DistT^2 + 1)^{sp_par/2})
Y \leftarrow rmnorm(1, rep(alpha, times = n), SIGMA) #generate the linear variable
theta <- c()
## wrapping step
for(i in 1:n) {
 theta[i] <- Y[i] %% (2*pi)
### Add plots of the simulated data
rose_diag(theta)
## use this values as references for the definition of initial values and priors
rho_sp.min <- 3/max(Dist)</pre>
rho_sp.max <- rho_sp.min+0.5</pre>
rho_t.min <- 3/max(DistT)</pre>
rho_t.max <- rho_t.min+0.5</pre>
val <- sample(1:n,round(n*0.2)) #validation set</pre>
set.seed(100)
mod <- WrapSpTi(</pre>
          = theta[-val],
          = coords[-val,],
 coords
 times
          = coordsT[-val],
 start = list("alpha"
                              = c(.79, .74),
                 "rho_sp"
                              = c(.33, .52),
                 "rho_t"
                             = c(.19, .43),
                 "sigma2"
                             = c(.49, .37),
                 "sep_par" = c(.47, .56),
                 "k"
                           = sample(0,length(theta[-val]), replace = TRUE)),
          = list("rho_sp"
 priors
                                = c(0.01,3/4), ### uniform prior on this interval
                  "rho_t"
                              = c(0.01,3/4), ### uniform prior on this interval
                  "sep_par" = c(1,1), ### beta prior
                  "sigma2" = c(5,5),## inverse gamma prior with mode=5/6
                  "alpha" = c(0,20) ## wrapped gaussian with large variance
 sd_prop = list( "sigma2" = 0.1, "rho_sp" = 0.1, "rho_t" = 0.1, "sep_par"= 0.1),
 iter = 7000,
 BurninThin
              = c(burnin = 3000, thin = 10),
 accept_ratio = 0.234,
 adapt_param = c(start = 1, end = 1000, exp = 0.5),
 n_{chains} = 2 ,
 parallel = FALSE,
 n_{cores} = 1
)
check <- ConvCheck(mod,startit = 1 ,thin = 1)</pre>
check$Rhat ## convergence has been reached
## when plotting chains remember that alpha is a circular variable
par(mfrow = c(3,2))
```

WrapSp 31

```
coda::traceplot(check$mcmc)
par(mfrow = c(1,1))

############## Prediction on the validation set
Krig <- WrapKrigSpTi(
   WrapSpTi_out = mod,
   coords_obs = coords[-val,],
   coords_nobs = coords[val,],
   times_obs = coordsT[-val],
   times_nobs = coordsT[val],
   x_obs = theta[-val]
)

### checking the prediction
Wrap_Ape <- APEcirc(theta[val], Krig$Prev_out)
Wrap_Crps <- CRPScirc(theta[val], Krig$Prev_out)</pre>
```

WrapSp

Samples from the Wrapped Normal spatial model

Description

The function WrapSp produces samples from the posterior distribution of the wrapped normal spatial model.

Usage

```
WrapSp(x = x, coords = coords, start = list(alpha = c(2, 1), rho =
  c(0.1, 0.5), sigma2 = c(0.1, 0.5), k = sample(0, length(x), replace =
  T)), priors = list(alpha = c(pi, 1, -10, 10), rho = c(8, 14), sigma2 =
  c()), sd_prop = list(sigma2 = 0.5, rho = 0.5), iter = 1000,
  BurninThin = c(burnin = 20, thin = 10), accept_ratio = 0.234,
  adapt_param = c(start = 1, end = 1e+07, exp = 0.9),
  corr_fun = "exponential", kappa_matern = 0.5, n_chains = 1,
  parallel = FALSE, n_cores = 1)
```

Arguments

x a vector of n circular data in $[0, 2\pi)$ If they are not in $[0, 2\pi)$, the function will tranform the data in the right interval

coords an nx2 matrix with the sites coordinates

start a list of 4 elements giving initial values for the model parameters. Each elements is a numeric vector with n_chains elements

- alpha the mean which value is in $[0, 2\pi)$.
- rho the spatial decay parameter
- sigma2 the process variance
- k the vector of length(x) winding numbers

32 WrapSp

priors a list of 3 elements to define priors for the model parameters: **alpha** a vector of 2 elements the mean and the variance of a Wrapped Gaussian distribution, default is mean π and variance 1, **rho** a vector of 2 elements defining the minimum and maximum of a uniform distribution, sigma2 a vector of 2 elements defining the shape and rate of an inverse-gamma distribution, sd_prop list of 3 elements. To run the MCMC for the rho and sigma2 parameters we use an adaptive metropolis and in sd.prop we build a list of initial guesses for these two parameters and the beta parameter number of iterations iter BurninThin a vector of 2 elements with the burnin and the chain thinning it is the desired acceptance ratio in the adaptive metropolis accept_ratio a vector of 3 elements giving the iteration number at which the adaptation must adapt_param start and end. The third element (exp) must be a number in (0,1) and it is a parameter ruling the speed of changes in the adaptation algorithm, it is recommended to set it close to 1, if it is too small non positive definite matrices may be generated and the program crashes. corr_fun characters, the name of the correlation function; currently implemented functions are c("exponential", "matern", "gaussian") numeric, the smoothness parameter of the Matern correlation function, default kappa_matern is kappa_matern = 0.5 (the exponential function) n_chains integer, the number of chains to be launched (default is 1, but we recommend to use at least 2 for model diagnostic) parallel logical, if the multiple chains must be lunched in parallel (you should install

Value

n_cores

It returns a list of n_chains lists each with elements

- alpha, rho, sigma2 vectors with the thinned chains,
- k a matrix with nrow = length(x) and ncol = the length of thinned chains

doParallel package). Default is FALSE

mentation. Default value is 1.

- corr_fun characters with the type of spatial correlation chosen.
- distribution characters, always "WrapSp"

Implementation Tips

To facilitate the estimations, the observations x are centered around pi, and the prior and starting value of alpha are changed accordingly. After the estimations, posterior samples of alpha are changed back to the original scale

integer, required if parallel=TRUE, the number of cores to be used in the imple-

WrapSp 33

References

G. Jona Lasinio, A. Gelfand, M. Jona-Lasinio, "Spatial analysis of wave direction data using wrapped Gaussian processes", The Annals of Applied Statistics 6 (2013), 1478-1498

See Also

WrapKrigSp for spatial interpolation, ProjSp for posterior sampling from the Projected Normal model and ProjKrigSp for spatial interpolation under the same model

```
library(CircSpaceTime)
## auxiliary function
rmnorm<-function(n = 1, mean = rep(0, d), varcov){
  d <- if (is.matrix(varcov))</pre>
    ncol(varcov)
  else 1
  z <- matrix(rnorm(n * d), n, d) %*% chol(varcov)</pre>
  y \leftarrow t(mean + t(z))
  return(y)
}
# Simulation with exponential spatial covariance function
####
set.seed(1)
n <- 20
coords <- cbind(runif(n,0,100), runif(n,0,100))
Dist <- as.matrix(dist(coords))</pre>
        <- 0.05
rho
sigma2 <- 0.3
alpha <-c(0.5)
       <- sigma2*exp(-rho*Dist)
Y <- rmnorm(1,rep(alpha,times=n), SIGMA)
theta <- c()
for(i in 1:n) {
  theta[i] <- Y[i]%%(2*pi)
rose_diag(theta)
#validation set
val <- sample(1:n,round(n*0.1))</pre>
set.seed(12345)
mod <- WrapSp(</pre>
     = theta[-val],
  coords = coords[-val,],
  start = list("alpha" = c(.36, 0.38),
                 "rho"
                        = c(0.041, 0.052),
                  "sigma2" = c(0.24, 0.32),
```

```
= rep(0,(n - length(val)))),
 priors = list("rho"
                         = c(0.04, 0.08), #few observations require to be more informative
                  "sigma2"
                             = c(2,1),
                  "alpha" = c(0,10)
 sd_prop = list( "sigma2" = 0.1, "rho" = 0.1),
         = 1000,
 BurninThin
               = c(burnin = 500, thin = 5),
 accept_ratio = 0.234,
 adapt_param = c(start = 40000, end = 45000, exp = 0.5),
 corr_fun = "exponential",
 kappa_matern = .5,
 parallel = FALSE,
 #With doParallel, bigger iter (normally around 1e6) and n_cores>=2 it is a lot faster
 n_{chains} = 2,
 n\_cores = 1
)
check <- ConvCheck(mod)</pre>
check$Rhat ## close to 1 means convergence has been reached
## graphical check
par(mfrow = c(3,1))
coda::traceplot(check$mcmc)
par(mfrow = c(1,1))
##### We move to the spatial interpolation see WrapKrigSp
```

WrapSpTi

Samples from the posterior distribution of the Wrapped Normal spatial temporal model

Description

The WrapSpTi function returns samples from the posterior distribution of the spatio-temporal Wrapped Gaussian Model

Usage

```
WrapSpTi(x = x, coords = coords, times, start = list(alpha = c(2, 1),
    rho_sp = c(0.1, 0.5), rho_t = c(0.1, 1), sep_par = c(0.01, 0.1), k =
    sample(0, length(x), replace = T)), priors = list(alpha = c(pi, 1, -10,
    10), rho_sp = c(8, 14), rho_t = c(1, 2), sep_par = c(0.001, 1), sigma2 =
    c()), sd_prop = list(rho_sp = 0.5, rho_t = 0.5, sep_par = 0.5, sigma2 =
    0.5), iter = 1000, BurninThin = c(burnin = 20, thin = 10),
    accept_ratio = 0.234, adapt_param = c(start = 1, end = 1e+07, exp =
    0.9), n_chains = 1, parallel = FALSE, n_cores = 1)
```

Arguments

x a vector of n circular data in $[0, 2\pi)$. If they are not in $[0, 2\pi)$, the function will transform the data into the right interval

coords an nx2 matrix with the sites coordinates times an n vector with the times of the observations x a list of 4 elements giving initial values for the model parameters. Each elements start is a vector with n chains elements • alpha the mean which value is in $[0, 2\pi)$ • rho_sp the spatial decay parameter, • rho_t the temporal decay parameter, • sigma2 the process variance, • sep_par the separation parameter, • k the vector of length(x) winding numbers a list of 5 elements to define priors for the model parameters: priors alpha a vector of 2 elements the mean and the variance of a Gaussian distribution, default is mean π and variance 1, **rho_sp** a vector of 2 elements defining the minimum and maximum of a uniform distribution, **rho_t** a vector of 2 elements defining the minimum and maximum of a uniform distribution, sep_par a vector of 2 elements defining the two parameters of a beta distribu**sigma2** a vector of 2 elements defining the shape and rate of an inverse-gamma distribution, list of 3 elements. To run the MCMC for the rho_sp and sigma2 parameters we sd_prop use an adaptive metropolis and in sd_prop we build a list of initial guesses for these two parameters and the beta parameter iter iter number of iterations BurninThin a vector of 2 elements with the burnin and the chain thinning accept_ratio it is the desired acceptance ratio in the adaptive metropolis adapt_param a vector of 3 elements giving the iteration number at which the adaptation must start and end. The third element (exp) must be a number in (0,1) and it is a parameter ruling the speed of changes in the adaptation algorithm, it is recommended to set it close to 1, if it is too small non positive definite matrices may

n_chains

integer, the number of chains to be launched (default is 1, but we recommend to

use at least 2 for model diagnostic)

parallel logical, if the multiple chains must be lunched in parallel (you should install

doParallel package). Default is FALSE

be generated and the program crashes.

n_cores integer, required if parallel=TRUE, the number of cores to be used in the imple-

mentation. Default value is 1.

Value

it returns a list of n_chains lists each with elements

alpha, rho_sp, rho_t, sep_par, sigma2 vectors with the thinned chains k a matrix with nrow = length(x) and ncol = the length of thinned chains distribution characters, always "WrapSpTi"

Implementation Tips

To facilitate the estimations, the observations x are centered around pi, and the prior and starting value of alpha are changed accordingly. After the estimations, posterior samples of alpha are changed back to the original scale

References

- G. Mastrantonio, G. Jona Lasinio, A. E. Gelfand, "Spatio-temporal circular models with non-separable covariance structure", TEST 25 (2016), 331–350.
- T. Gneiting, "Nonseparable, Stationary Covariance Functions for Space-Time Data", JASA 97 (2002), 590-600

See Also

WrapKrigSpTi for spatio-temporal prediction, ProjSpTi to sample from the posterior distribution of the spatio-temporal Projected Normal model and ProjKrigSpTi for spatio-temporal prediction under the same model

Other spatio-temporal models: ProjSpTi

```
library(CircSpaceTime)
## functions
rmnorm <- function(n = 1, mean = rep(0, d), varcov){
 d <- if (is.matrix(varcov))</pre>
   ncol(varcov)
 else 1
 z <- matrix(rnorm(n * d), n, d) %*% chol(varcov)</pre>
 y \leftarrow t(mean + t(z))
 return(y)
## Simulation
set.seed(1)
n <- 20
### simulate coordinates from a unifrom distribution
coords < cbind(runif(n,0,100), runif(n,0,100)) #spatial coordinates
coordsT <- sort(runif(n,0,100)) #time coordinates (ordered)</pre>
Dist <- as.matrix(dist(coords))</pre>
DistT <- as.matrix(dist(coordsT))</pre>
rho
       <- 0.05 #spatial decay
       <- 0.01 #temporal decay
sep_par <- 0.5 #separability parameter</pre>
sigma2 <- 0.3 # variance of the process
alpha <-c(0.5)
#Gneiting covariance
```

```
SIGMA \leftarrow sigma2 * (rhoT * DistT^2 + 1)^(-1) * exp(-rho * Dist/(rhoT * DistT^2 + 1)^(sep_par/2))
Y \leftarrow rmnorm(1, rep(alpha, times = n), SIGMA) #generate the linear variable
theta <- c()
## wrapping step
for(i in 1:n) {
 theta[i] <- Y[i] %% (2*pi)
### Add plots of the simulated data
rose_diag(theta)
## use this values as references for the definition of initial values and priors
rho_sp.min <- 3/max(Dist)</pre>
rho_sp.max <- rho_sp.min+0.5</pre>
rho_t.min <- 3/max(DistT)</pre>
rho_t.max <- rho_t.min+0.5</pre>
val <- sample(1:n,round(n*0.2)) #validation set</pre>
set.seed(100)
mod <- WrapSpTi(</pre>
 Χ
          = theta[-val],
          = coords[-val,],
 coords
         = coordsT[-val],
 times
 start = list("alpha"
                              = c(.79, .74),
                 "rho_sp"
                              = c(.33,.52),
                 "rho_t"
                             = c(.19, .43),
                 "sigma2"
                             = c(.49, .37),
                 "sep_par" = c(.47, .56),
                 "k"
                       = sample(0,length(theta[-val]), replace = TRUE)),
          = list("rho_sp"
                               = c(0.01,3/4), ### uniform prior on this interval
 priors
                  "rho_t"
                               = c(0.01,3/4), ### uniform prior on this interval
                  "sep_par" = c(1,1), ### beta prior
                  "sigma2"
                             = c(5,5),## inverse gamma prior with mode=5/6
                  "alpha" = c(0,20) ## wrapped gaussian with large variance
 sd_prop = list( "sigma2" = 0.1, "rho_sp" = 0.1, "rho_t" = 0.1, "sep_par"= 0.1),
          = 7000,
 iter
              = c(burnin = 3000, thin = 10),
 BurninThin
 accept_ratio = 0.234,
 adapt_param = c(start = 1, end = 1000, exp = 0.5),
 n_{chains} = 2 ,
 parallel = FALSE,
 n\_cores = 1
)
check <- ConvCheck(mod,startit = 1 ,thin = 1)</pre>
check$Rhat ## convergence has been reached
## when plotting chains remember that alpha is a circular variable
par(mfrow = c(3,2))
coda::traceplot(check$mcmc)
par(mfrow = c(1,1))
#### move to the prediction step with WrapKrigSpTi
```

Index

rose_diag, 25

```
* convergence check indices
                                                     WrapKrigSp, 2, 3, 6, 7, 9, 12, 19, 26, 33
    ConvCheck, 6
                                                     WrapKrigSpTi, 2, 3, 6, 7, 9, 15, 23, 28, 36
* datasets
                                                     WrapSp, 6, 12, 19, 26, 31
    april, 5
                                                     WrapSpTi, 6, 15, 23, 28, 29, 34
    may, 10
* model performance indices
    APEcirc, 2
    CRPScirc, 8
* spatial estimations
    WrapSp, 31
* spatial interpolations
    ProjKrigSp, 11
    WrapKrigSp, 26
* spatial models
    ProjSp, 17
* spatio-temporal estimations
    WrapKrigSpTi, 28
* spatio-temporal models
    ProjSpTi, 21
    WrapSpTi, 34
* spato-temporal interpolations
    ProjKrigSpTi, 14
APEcirc, 2, 9
april, 5
CircSpaceTime, 6
CircSpaceTime-package (CircSpaceTime), 6
ConvCheck, 6
CRPScirc, 3, 8
gelman.diag, 6
may, 10
ProjKrigSp, 2, 3, 6, 7, 9, 11, 19, 26, 33
ProjKrigSpTi, 2, 3, 6, 7, 9, 14, 23, 29, 36
ProjSp, 6, 11, 12, 17, 26, 33
ProjSpTi, 6, 14, 15, 21, 29, 36
```