Package 'spNNGP'

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Title Spatial Regression Models for Large Datasets using Nearest Neighbor Gaussian Processes		
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Maintainer Andrew Finley <finleya@msu.edu></finleya@msu.edu>		
Author Andrew Finley [aut, cre], Abhirup Datta [aut], Sudipto Banerjee [aut], Alexander Mckim [ctb]		
Depends R (> 2.10.0), coda, Formula, RANN		
Description Fits Gaussian univariate Bayesian spatial regression models for large datasets using Nea est Neighbor Gaussian Processes (NNGP) detailed in Datta, Banerjee, Finley, and Gelfand (2016) <doi:10.1080 01621459.2015.1044091=""> and Finley, Datta, Cook, Morton, Andersen, and Banerjee (2017) <arxiv:1702.00434v2>.</arxiv:1702.00434v2></doi:10.1080>		
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R topics documented:		
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СНМ	Canopy Height Model from NASA Goddard's LiDAR Hyperspectral and Thermal (G-LiHT)
	= (& =)

Description

Canopy Height Model (CHM) from NASA Goddard's LiDAR Hyperspectral and Thermal (G-LiHT; Cook et al. 2013) Airborne Imager over a subset of Harvard Forest Simes Tract, MA, collected in Summer 2012.

The CHM matrix columns are longitude, latitude, and canopy height (m) from ground for 17,23,137 observations. Longitude and latitude are in UTM Zone 18 (proj4string "+proj=utm +zone=18 +da-tum=WGS84 +units=m +no_defs +ellps=WGS84 +towgs84=0,0,0").

Usage

data(CHM)

Format

A matrix containing 17,23,137 rows and 3 columns named longitude, latitude, and CHM.

Source

Data were downloaded from https://gliht.gsfc.nasa.gov with metadata available at ftp://fusionftp.gsfc.nasa.gov/G-LiHT/Simes_Jun2012/metadata/Simes_Jun2012_metadata.pdf.

References

Cook, B.D., L.W. Corp, R.F. Nelson, E.M. Middleton, D.C. Morton, J.T. McCorkel, J.G. Masek, K.J. Ranson, and V. Ly. (2013) NASA Goddard's Lidar, Hyperspectral and Thermal (G-LiHT) airborne imager. Remote Sensing 5:4045-4066.

spConjNNGP	Function for fitting univariate Bayesian conjugate spatial regression models
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Description

The function spConjNNGP fits Gaussian univariate Bayesian conjugate spatial regression models using Nearest Neighbor Gaussian Processes (NNGP).

Usage

Arguments

formula a symbolic description of the regression model to be fit. See example below.

data an optional data frame containing the variables in the model. If not found in data,

the variables are taken from environment (formula), typically the environment

from which spConjNNGP is called.

coords an $n \times 2$ matrix of the observation coordinates in R^2 (e.g., easting and northing).

n.neighbors number of neighbors used in the NNGP.

theta.alpha a vector or matrix of parameter values for phi, nu, and alpha, where $\alpha = \tau^2/\sigma^2$

and nu is only required if cov.model="matern". A vector is passed to run the model using one set of parameters. The vector elements must be named and hold values for phi, nu, and alpha. If a matrix is passed, columns must be named and hold values for phi, alpha, and nu. Each row in the matrix defines a set of

parameters for which the model will be run.

sigma.sq.IG a vector of length two that holds the hyperparameters, *shape* and *scale* respec-

tively, for the inverse-Gamma prior on σ^2 .

cov.model a quoted keyword that specifies the covariance function used to model the spatial

dependence structure among the observations. Supported covariance model key words are: "exponential", "matern", "spherical", and "gaussian". See

below for details.

k. fold specifies the number of k folds for cross-validation. If theta. alpha is a vector

then cross-validation is not performed and k-fold and score.rule are ignored. In k-fold cross-validation, the data specified in model is randomly partitioned into k equal sized subsamples. Of the k subsamples, k-1 subsamples are used to fit the model and the remaining k samples are used for prediction. The cross-validation process is repeated k times (the folds). Root mean squared prediction error (RMSPE) and continuous ranked probability score (CRPS; Gneiting and Raftery, 2007) rules are averaged over the k fold prediction results and reported for the parameter sets defined by theta.alpha. The parameter set that yields the *best* performance based on the scoring rule defined by score.rule is used fit the final model that uses all the data and make predictions if X.0 and coords.0 are specified. Results from the k-fold cross-validation are returned in

the k. fold. scores matrix.

a quoted keyword "rmspe" or "crps" that specifies the scoring rule used to select the *best* parameter set, see argument definition for k. fold for more details.

X.0 the design matrix for prediction locations. An intercept should be provided in

the first column if one is specified in model.

coords.0 the spatial coordinates corresponding to X.0.

n.omp.threads a positive integer indicating the number of threads to use for SMP parallel pro-

cessing. The package must be compiled for OpenMP support. For most Intel-based machines, we recommend setting n.omp.threads up to the number of

hyperthreaded cores.

search.type a quoted keyword that specifies type of nearest neighbor search algorithm. Sup-

ported method key words are: "tree" and "brute" both will yield the same

solution but "tree" should be much faster.

return.neighbors

if TRUE, a list containing the indices for each location's nearest neighbors will be returned along with ordered data used to fit a NNGP model. This argument

should typically be FALSE. See n. indx below for more details.

verbose if TRUE, model specification and progress is printed to the screen. Otherwise,

nothing is printed to the screen.

... currently no additional arguments.

Value

An object of class cNNGP, which is a list comprising:

theta.alpha the input theta.alpha vector, or best (according to the selected scoring rule) set

of parameters in the theta.alpha matrix. All subsequent parameter estimates

are based on this parameter set.

beta. hat a matrix of regression coefficient estimates corresponding to the returned theta. alpha.

beta.var beta.hat variance-covariance matrix.

sigma.sq.hat estimate of σ^2 corresponding to the returned theta.alpha.

sigma.sq.var sigma.sq.hat variance.

k.fold.scores results from the k-fold cross-validation if theta.alpha is a matrix.

y.0.hat prediction if X.0 and coords.0 are specified.

y.0.var.hat y.0.hat variance.

n.neighbors number of neighbors used in the NNGP.

n.indx if return.neighbors=TRUE then n.indx will be a list of length n. The i-th

element in the list corresponds to the i-th row in coords.ord matrix and the

elements are the nearest neighbor indices for the given location.

ord if return.neighbors=TRUE the vector ord=order(coords[,1]), which is the

vector of indices used to order data necessary for fitting the NNGP model, is

returned.

coords.ord if return.neighbors=TRUE then coords.ord = coords[ord,] is returned.

y.ord if return.neighbors=TRUE then y.ord = y[ord] is returned.

X.ord if return.neighbors=TRUE then X.ord = X[ord,,drop=FALSE] is returned.

run.time execution time for building the nearest neighbor index and parameter estimation

reported using proc.time().

Author(s)

```
Andrew O. Finley <finleya@msu.edu>,
Abhirup Datta <abhidatta@jhu.edu>,
Sudipto Banerjee <sudipto@ucla.edu>
```

References

Datta, A., S. Banerjee, A.O. Finley, and A.E. Gelfand. (2016) Hierarchical Nearest-Neighbor Gaussian process models for large geostatistical datasets. Journal of the American Statistical Association, 111:800-812.

Finley, A.O., A. Datta, B.C. Cook, D.C. Morton, H.E. Andersen, and S. Banerjee (2017) Applying Nearest Neighbor Gaussian Processes to massive spatial data sets: Forest canopy height prediction across Tanana Valley Alaska, https://arxiv.org/abs/1702.00434v2.

Gneiting, T and A.E. Raftery. (2007) Strictly proper scoring rules, prediction, and estimation. Journal of the American Statistical Association, 102:359-378.

Examples

```
rmvn <- function(n, mu=0, V = matrix(1)){</pre>
  p <- length(mu)</pre>
  if(any(is.na(match(dim(V),p))))
    stop("Dimension problem!")
  D <- chol(V)
  t(matrix(rnorm(n*p), ncol=p)%*%D + rep(mu,rep(n,p)))
##Make some data
set.seed(1)
n <- 2000
coords <- cbind(runif(n,0,1), runif(n,0,1))
x <- cbind(1, rnorm(n))</pre>
B \leftarrow as.matrix(c(1,5))
sigma.sq <- 5
tau.sq <- 1
phi <- 3/0.5
D <- as.matrix(dist(coords))</pre>
R <- exp(-phi*D)
w \leftarrow rmvn(1, rep(0,n), sigma.sq*R)
y \leftarrow rnorm(n, x\%*B + w, sqrt(tau.sq))
ho <- sample(1:n, 1000)
y.ho \leftarrow y[ho]
x.ho <- x[ho,,drop=FALSE]</pre>
w.ho \leftarrow w[ho]
```

```
coords.ho <- coords[ho,]</pre>
y \leftarrow y[-ho]
x <- x[-ho,,drop=FALSE]</pre>
w <- w[-ho,,drop=FALSE]</pre>
coords <- coords[-ho,]</pre>
##Fit a Conjugate NNGP model and predict for the holdout
sigma.sq.IG <- c(2, sigma.sq)</pre>
cov.model <- "exponential"</pre>
g <- 10
theta.alpha <- cbind(seq(phi,30,length.out=g), seq(tau.sq/sigma.sq,5,length.out=g))</pre>
colnames(theta.alpha) <- c("phi", "alpha")</pre>
##one thread
m.c \leftarrow spConjNNGP(y^x-1, coords=coords, n.neighbors = 10,
                   X.0 = x.ho, coords.0 = coords.ho,
                   k.fold = 5, score.rule = "crps",
                   n.omp.threads = 1,
                theta.alpha = theta.alpha, sigma.sq.IG = sigma.sq.IG, cov.model = cov.model)
m.c$beta.hat
m.c$theta.alpha.sigmaSq
m.c$k.fold.scores
##two threads
m.c \leftarrow spConjNNGP(y^x-1, coords=coords, n.neighbors = 10,
                   X.0 = x.ho, coords.0 = coords.ho,
                   k.fold = 5, score.rule = "crps",
                   n.omp.threads = 2,
                theta.alpha = theta.alpha, sigma.sq.IG = sigma.sq.IG, cov.model = cov.model)
m.c$beta.hat
m.c$sigmaSq.hat
m.c$k.fold.scores
```

spNNGP

Function for fitting univariate Bayesian spatial regression models

Description

The function spNNGP fits Gaussian univariate Bayesian spatial regression models using Nearest Neighbor Gaussian Processes (NNGP).

Usage

```
spNNGP(formula, data = parent.frame(), coords, method = "response", n.neighbors = 15,
    starting, tuning, priors, cov.model = "exponential",
    n.samples, n.omp.threads = 1, search.type = "tree",
    return.neighbors = FALSE, verbose = TRUE, n.report = 100, ...)
```

Arguments

formula a symbolic description of the regression model to be fit. See example below.

data an optional data frame containing the variables in the model. If not found in data,

the variables are taken from environment (formula), typically the environment

from which spNNGP is called.

coords an $n \times 2$ matrix of the observation coordinates in R^2 (e.g., easting and northing).

method a quoted keyword that specifies the NNGP sampling algorithm. Supported

method keywords are: "response" and "sequential". When n is large, e.g., greater than 100k, the "response" algorithm should be faster. In general, unless estimates of spatial random effects are needed, the "response" algorithm

should be used. See below for details.

n.neighbors number of neighbors used in the NNGP.

starting a list with each tag corresponding to a parameter name. Valid tags are beta,

sigma.sq, tau.sq, phi, and nu. nu is only specified if cov.model="matern".

The value portion of each tag is the parameter's startingvalue.

tuning a list with each tag corresponding to a parameter name. Valid tags are sigma. sq,

tau.sq, phi, and nu. If method="sequential" then only phi and nu need to be specified. The value portion of each tag defines the variance of the Metropolis

sampler Normal proposal distribution.

priors a list with each tag corresponding to a parameter name. Valid tags are sigma.sq.ig,

tau.sq.ig, phi.unif, and nu.unif. Variance parameters, simga.sq and tau.sq, are assumed to follow an inverse-Gamma distribution, whereas the spatial decay phi and smoothness nu parameters are assumed to follow Uniform distributions. The hyperparameters of the inverse-Gamma are passed as a vector of length two, with the first and second elements corresponding to the *shape* and *scale*, respectively. The hyperparameters of the Uniform are also passed as a vector of length two with the first and second elements corresponding to the lower and upper

support, respectively.

cov.model a quoted keyword that specifies the covariance function used to model the spatial

dependence structure among the observations. Supported covariance model key words are: "exponential", "matern", "spherical", and "gaussian". See

below for details.

n. samples the number of posterior samples to collect.

n.omp.threads a positive integer indicating the number of threads to use for SMP parallel pro-

cessing. The package must be compiled for OpenMP support. For most Intelbased machines, we recommend setting n.omp.threads up to the number of

hyperthreaded cores.

verbose if TRUE, model specification and progress of the sampler is printed to the screen.

Otherwise, nothing is printed to the screen.

search.type a quoted keyword that specifies type of nearest neighbor search algorithm. Sup-

ported method key words are: "tree" and "brute" both will yield the same

solution but "tree" should be much faster.

return.neighbors

if TRUE, a list containing the indices for each location's nearest neighbors will be returned along with ordered data used to fit a NNGP model. This argument

should typically be FALSE. See n. indx below for more details.

n.report the interval to report Metropolis sampler acceptance and MCMC progress.

... currently no additional arguments.

Details

Model parameters can be fixed at their starting values by setting their tuning values to zero.

The *no nugget* model is specified by setting tau. sq to zero in the starting and tuning lists.

Value

An object of class rNNGP or sNNGP depending on the method, which is a list comprising:

p.beta.samples a coda object of posterior samples for the regression coefficients.

p.theta.samples

a coda object of posterior samples for covariance parameters.

p.w.samples is a matrix of posterior samples for the spatial random effects, where rows cor-

respond to locations in coords and columns hold the n. samples posterior sam-

ples. This is only returned if method="sequential".

n.indx if return.neighbors=TRUE then n.indx will be a list of length n. The i-th

element in the list corresponds to the i-th row in coords.ord matrix and the

elements are the nearest neighbor indices for the given location.

ord the vector order (coords[,1]), which is the vector of indices used to order data

necessary for fitting the NNGP model.

coords.ord the matrix coords[ord,].

y.ord the vector y[ord].

X. ord the matrix X[ord, ,drop=FALSE].

run.time execution time for building the nearest neighbor index and MCMC sampler re-

ported using proc. time().

The return object will include additional objects used for subsequent prediction and/or model fit evaluation.

Author(s)

Andrew O. Finley <finleya@msu.edu>, Abhirup Datta <abhidatta@jhu.edu>, Sudipto Banerjee <sudipto@ucla.edu>

References

Datta, A., S. Banerjee, A.O. Finley, and A.E. Gelfand. (2016) Hierarchical Nearest-Neighbor Gaussian process models for large geostatistical datasets. Journal of the American Statistical Association, 111:800-812

Finley, A.O., A. Datta, B.C. Cook, D.C. Morton, H.E. Andersen, and S. Banerjee (2017) Applying Nearest Neighbor Gaussian Processes to massive spatial data sets: Forest canopy height prediction across Tanana Valley Alaska, https://arxiv.org/abs/1702.00434v2.

Examples

```
rmvn <- function(n, mu=0, V = matrix(1)){</pre>
  p <- length(mu)</pre>
  if(any(is.na(match(dim(V),p))))
    stop("Dimension problem!")
  D <- chol(V)
  t(matrix(rnorm(n*p), ncol=p)%*%D + rep(mu,rep(n,p)))
}
##Make some data
set.seed(1)
n <- 100
coords <- cbind(runif(n,0,1), runif(n,0,1))</pre>
x <- cbind(1, rnorm(n))</pre>
B \leftarrow as.matrix(c(1,5))
sigma.sq <- 5
tau.sq <- 1
phi <- 3/0.5
D <- as.matrix(dist(coords))</pre>
R <- exp(-phi*D)
w \leftarrow rmvn(1, rep(0,n), sigma.sq*R)
y \leftarrow rnorm(n, x\%*B + w, sqrt(tau.sq))
##Fit a Response and Sequential NNGP model
n.samples < -500
starting <- list("phi"=phi, "sigma.sq"=5, "tau.sq"=1)</pre>
tuning <- list("phi"=0.1, "sigma.sq"=0.1, "tau.sq"=0.1)</pre>
priors <- list("phi.Unif"=c(3/1, 3/0.01), "sigma.sq.IG"=c(2, 5), "tau.sq.IG"=c(2, 1))
cov.model <- "exponential"</pre>
m.s <- spNNGP(y~x-1, coords=coords, starting=starting, method="sequential", n.neighbors=10,
               tuning=tuning, priors=priors, cov.model=cov.model,
               n.samples=n.samples, n.omp.threads=2)
```

spPredict

spPredict

Function for prediction at new locations using spNNGP models.

Description

The function spPredict collects posterior predictive samples for a set of new locations given a spNNGP object.

Usage

Arguments

sp.obj	an object returned by spNNGP.
X.0	the design matrix for prediction locations. An intercept should be provided in the first column if one is specified in sp.obj model.
coords.0	the spatial coordinates corresponding to X. Ø.
start	specifies the first sample included in the composition sampling.
end	specifies the last sample included in the composition. The default is to use all posterior samples in sp.obj.
thin	a sample thinning factor. The default of 1 considers all samples between start and end. For example, if thin = 10 then 1 in 10 samples are considered between start and end.
n.omp.threads	a positive integer indicating the number of threads to use for SMP parallel processing. The package must be compiled for OpenMP support. For most Intelbased machines, we recommend setting n.omp.threads up to the number of hyperthreaded cores.
verbose	if TRUE, model specification and progress of the sampler is printed to the screen. Otherwise, nothing is printed to the screen.
n.report	the interval to report sampling progress.
	currently no additional arguments.

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Value

A list comprising:

p.y.0	a matrix that holds the response variable posterior predictive samples where rows are locations corresponding to coords.0 and columns are samples.
p.w.0	a matrix that holds the random effect posterior predictive samples where rows are locations corresponding to coords.0 and columns are samples. This is only returned if spNNGP method = "sequential".
run.time	execution time reported using proc.time().

Author(s)

```
Andrew O. Finley <finleya@msu.edu>,
Abhirup Datta <abhidatta@jhu.edu>,
Sudipto Banerjee <sudipto@ucla.edu>
```

References

Datta, A., S. Banerjee, A.O. Finley, and A.E. Gelfand. (2016) Hierarchical Nearest-Neighbor Gaussian process models for large geostatistical datasets. Journal of the American Statistical Association, 111:800-812.

Finley, A.O., A. Datta, B.C. Cook, D.C. Morton, H.E. Andersen, and S. Banerjee (2017) Applying Nearest Neighbor Gaussian Processes to massive spatial data sets: Forest canopy height prediction across Tanana Valley Alaska, https://arxiv.org/abs/1702.00434v2.

Examples

```
rmvn <- function(n, mu=0, V = matrix(1)){</pre>
  p <- length(mu)</pre>
  if(any(is.na(match(dim(V),p))))
    stop("Dimension problem!")
  D <- chol(V)
  t(matrix(rnorm(n*p), ncol=p)%*%D + rep(mu,rep(n,p)))
}
##Make some data
set.seed(1)
n <- 100
coords <- cbind(runif(n,0,1), runif(n,0,1))
x <- cbind(1, rnorm(n))</pre>
B \leftarrow as.matrix(c(1,5))
sigma.sq <- 5
tau.sq <- 1
phi <- 3/0.5
D <- as.matrix(dist(coords))</pre>
```

spPredict spPredict

```
R <- exp(-phi*D)</pre>
w <- rmvn(1, rep(0,n), sigma.sq*R)</pre>
y \leftarrow rnorm(n, x%*B + w, sqrt(tau.sq))
ho <- sample(1:n, 50)
y.ho \leftarrow y[ho]
x.ho <- x[ho,,drop=FALSE]</pre>
w.ho <- w[ho]
coords.ho <- coords[ho,]</pre>
y \leftarrow y[-ho]
x <- x[-ho,,drop=FALSE]</pre>
w <- w[-ho,,drop=FALSE]</pre>
coords <- coords[-ho,]</pre>
##Fit a Response and Sequential NNGP model
n.samples <- 500
starting <- list("phi"=phi, "sigma.sq"=5, "tau.sq"=1)</pre>
tuning <- list("phi"=0.1, "sigma.sq"=0.1, "tau.sq"=0.1)</pre>
priors <- list("phi.Unif"=c(3/1, 3/0.01), "sigma.sq.IG"=c(2, 5), "tau.sq.IG"=c(2, 1))</pre>
cov.model <- "exponential"</pre>
n.report <- 500
##Predict for holdout set using both models
\verb|m.s| <- spNNGP(y^x-1, coords=coords, starting=starting, method="sequential", n.neighbors=10, method=10, me
                                     tuning=tuning, priors=priors, cov.model=cov.model,
                                    n.samples=n.samples, n.omp.threads=2, n.report=n.report)
m.r <- spNNGP(y~x-1, coords=coords, starting=starting, method="response", n.neighbors=10,</pre>
                                     tuning=tuning, priors=priors, cov.model=cov.model,
                                    n.samples=n.samples, n.omp.threads=2, n.report=n.report)
##Prediction for holdout data
p.s <- spPredict(m.s, X.0 = x.ho, coords.0 = coords.ho, n.omp.threads=2)
plot(apply(p.s$p.w.0, 1, mean), w.ho)
plot(apply(p.s$p.y.0, 1, mean), y.ho)
p.r \leftarrow spPredict(m.r, X.0 = x.ho, coords.0 = coords.ho, n.omp.threads=2)
plot(apply(p.r$p.y.0, 1, mean), y.ho)
```

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