Package ‘spmodel’

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Title Spatial Statistical Modeling and Prediction

Version 0.6.0

Description Fit, summarize, and predict for a variety of spatial statistical models applied to point-referenced and areal (lattice) data. Parameters are estimated using various methods. Additional modeling features include anisotropy, non-spatial random effects, partition factors, big data approaches, and more. Model-fit statistics are used to summarize, visualize, and compare models. Predictions at unobserved locations are readily obtainable. For additional details, see Dumelle et al. (2023) <doi:10.1371/journal.pone.0282524>.

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AIC.spmodel

Compute AIC and AICc of fitted model objects

Description

Compute AIC and AICc for one or several fitted model objects for which a log-likelihood value can be obtained.

Usage

```r
## S3 method for class 'splm'
AIC(object, ..., k = 2)

## S3 method for class 'spautor'
AIC(object, ..., k = 2)

## S3 method for class 'spglm'
AIC(object, ..., k = 2)

## S3 method for class 'spgautor'
AIC(object, ..., k = 2)

AICc(object, ..., k = 2)

## S3 method for class 'splm'
AICc(object, ..., k = 2)

## S3 method for class 'spautor'
AICc(object, ..., k = 2)

## S3 method for class 'spglm'
AICc(object, ..., k = 2)

## S3 method for class 'spgautor'
AICc(object, ..., k = 2)
```
Arguments

- **object**: A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()` where `estmethod` is "ml" or "reml".
- **...**: Optionally more fitted model objects.
- **k**: The penalty parameter, taken to be 2. Currently not allowed to differ from 2 (needed for generic consistency).

Details

When comparing models fit by maximum or restricted maximum likelihood, the smaller the AIC or AICc, the better the fit. The AICc contains a correction to AIC for small sample sizes. The theory of AIC and AICc requires that the log-likelihood has been maximized, and hence, no AIC or AICc methods exist for models where `estmethod` is not "ml" or "reml". Additionally, AIC and AICc comparisons between "ml" and "reml" models are meaningless – comparisons should only be made within a set of models estimated using "ml" or a set of models estimated using "reml". AIC and AICc comparisons for "reml" must use the same fixed effects. To vary the covariance parameters and fixed effects simultaneously, use "ml".

Hoeting et al. (2006) defines that spatial AIC as $-2\log\text{lik} + 2(estparams)$ and the spatial AICc as $-2\log\text{lik} + 2n(estparams)/(n - estparams - 1)$, where $n$ is the sample size and `estparams` is the number of estimated parameters. For "ml", `estparams` is the number of estimated covariance parameters plus the number of estimated fixed effects. For "reml", `estparams` is the number of estimated covariance parameters.

Value

- If just one object is provided, a numeric value with the corresponding AIC or AICc.
- If multiple objects are provided, a data.frame with rows corresponding to the objects and columns representing the number of parameters estimated (df) and the AIC or AICc.

Examples

```r
spmod <- splm(z ~ water + tarp,
  data = caribou,
  spcov_type = "exponential", xcoord = x, ycoord = y
)
AIC(spmod)
AICc(spmod)
```

**Description**

Compute analysis of variance and likelihood ratio tests of fitted model objects.
Usage

```r
## S3 method for class 'splm'
anova(object, ..., test = TRUE, Terms, L)

## S3 method for class 'spautor'
anova(object, ..., test = TRUE, Terms, L)

## S3 method for class 'spglm'
anova(object, ..., test = TRUE, Terms, L)

## S3 method for class 'spgautor'
anova(object, ..., test = TRUE, Terms, L)

## S3 method for class 'anova.splm'
tidy(x, ...)

## S3 method for class 'anova.spautor'
tidy(x, ...)

## S3 method for class 'anova.spglm'
tidy(x, ...)

## S3 method for class 'anova.spgautor'
tidy(x, ...)
```

Arguments

- **object**: A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`.
- **...**: An additional fitted model object.
- **test**: A logical value indicating whether p-values from asymptotic Chi-squared hypothesis tests should be returned. Defaults to `TRUE`.
- **Terms**: An optional character or integer vector that specifies terms in the model used to jointly compute test statistics and p-values (if `test = TRUE`) against a null hypothesis of zero. Terms is only used when a single fitted model object is passed to the function. If `Terms` is a character vector, it should contain the names of the fixed effect terms. If `Terms` is an integer vector, it should correspond to the order (starting at one) of the names of the fixed effect terms. The easiest way to obtain the names of all possible terms is to run `tidy(anova(object))$effects` (the integer representation matches the positions of this vector).
- **L**: An optional numeric matrix or list specifying linear combinations of the coefficients in the model used to compute test statistics and p-values (if `test = TRUE`) for coefficient constraints corresponding to a null hypothesis of zero. `L` is only used when a single fitted model object is passed to the function. If `L` is a numeric matrix, its rows indicate coefficient constraints and its columns represent coefficients. Then a single hypothesis test is conducted against a null hypothesis of zero. If `L` is a list, each list element is a numeric matrix specified as above.
separate hypothesis tests are conducted. The easiest way to obtain all possible coefficients is to run `tidy(object)$term`.

An object from `anova(object)`.

**Details**

When one fitted model object is present, `anova()` performs a general linear hypothesis test corresponding to some hypothesis specified by a matrix of constraints. If `Terms` and `L` are not specified, each model term is tested against zero (which correspond to type III or marginal hypothesis tests from classical ANOVA). If `Terms` is specified and `L` is not specified, all terms are tested jointly against zero. When `L` is specified, the linear combinations of terms specified by `L` are jointly tested against zero.

When two fitted model objects are present, one must be a "reduced" model nested in a "full" model. Then `anova()` performs a likelihood ratio test.

**Value**

When one fitted model object is present, `anova()` returns a data frame with degrees of freedom (Df), test statistics (Chi2), and p-values (Pr(>Chi2) if test = TRUE) corresponding to asymptotic Chi-squared hypothesis tests for each model term.

When two fitted model objects are present, `anova()` returns a data frame with the difference in degrees of freedom between the full and reduced model (Df), a test statistic (Chi2), and a p-value corresponding to the likelihood ratio test (Pr(>Chi2) if test = TRUE).

Whether one or two fitted model objects are provided, `tidy()` can be used to obtain tidy tibbles of the `anova(object)` output.

**Examples**

```
# one-model anova
spmod <- splm(z ~ water + tarp,
  data = caribou,
  spcov_type = "exponential", xcoord = x, ycoord = y
)
anova(spmod)
tidy(anova(spmod))
# see terms
tidy(anova(spmod))$effects
tidy(anova(spmod, Terms = c("water", "tarp")))
# same as
tidy(anova(spmod, Terms = c(2, 3)))
# likelihood ratio test
lmod <- splm(z ~ water + tarp,
  data = caribou,
  spcov_type = "none"
)
tidy(anova(spmod, lmod))
```
Description

Augment accepts a fitted model object and a data set and adds information about each observation in the data set. New columns always begin with a . prefix to avoid overwriting columns in the original data set.

Augment behaves differently depending on whether the original data or new data requires augmenting. Typically, when augmenting the original data, only the fitted model object is specified, and when augmenting new data, the fitted model object and newdata is specified. When augmenting the original data, diagnostic statistics are augmented to each row in the data set. When augmenting new data, predictions and optional intervals or standard errors are augmented to each row in the new data set.

Usage

```r
## S3 method for class 'spml'
augment(
  x,
  drop = TRUE,
  newdata = NULL,
  se_fit = FALSE,
  interval = c("none", "confidence", "prediction"),
  level = 0.95,
  local,
  ...
)
```

```r
## S3 method for class 'spautor'
augment(
  x,
  drop = TRUE,
  newdata = NULL,
  se_fit = FALSE,
  interval = c("none", "confidence", "prediction"),
  level = 0.95,
  local,
  ...
)
```

```r
## S3 method for class 'spglm'
augment(
  x,
  drop = TRUE,
  newdata = NULL,
```
Arguments

x A fitted model object from `splm()` or `spgautor()`.
drop A logical indicating whether to drop extra variables in the fitted model object `x` when augmenting. The default for `drop` is `TRUE`. `drop` is ignored if augmenting `newdata`.
newdata A data frame or tibble containing observations requiring prediction. All of the original explanatory variables used to create the fitted model object `x` must be present in `newdata`. Defaults to `NULL`, which indicates that nothing has been passed to `newdata`.
se_fit Logical indicating whether or not a `se.fit` column should be added to augmented output. Passed to `predict()` and defaults to `FALSE`.
interval Character indicating the type of confidence interval columns to add to the augmented `newdata` output. Passed to `predict()` and defaults to "none".
level Tolerance/confidence level. The default is 0.95.
local A list or logical. If a list, specific list elements described in `predict.spmodel()` control the big data approximation behavior. If a logical, `TRUE` chooses default list elements for the list version of `local` as specified in `predict.spmodel()`. Defaults to `FALSE`, which performs exact computations.
... Other arguments. Not used (needed for generic consistency).
type The scale (response or link) of predictions obtained using `spglm()` or `spgautor` objects.
newdata_size  The size value for each observation in newdata used when predicting for the binomial family.
var_correct  A logical indicating whether to return the corrected prediction variances when predicting via models fit using spglm() or spgautor(). The default is TRUE.

Details

augment() returns a tibble with the same class as data. That is, if data is an sf object, then the augmented object (obtained via augment(x)) will be an sf object as well. When augmenting newdata, the augmented object has the same class as data.

Missing response values from the original data can be augmented as if they were a newdata object by providing x$newdata to the newdata argument (where x is the name of the fitted model object). This is the only way to compute predictions for spautor() and spgautor() fitted model objects.

Value

When augmenting the original data set, a tibble with additional columns

- .fitted  Fitted value
- .resid  Response residual (the difference between observed and fitted values)
- .hat  Leverage (diagonal of the hat matrix)
- .cooksd  Cook’s distance
- .std.resid  Standardized residuals
- .se.fit  Standard error of the fitted value.

When augmenting a new data set, a tibble with additional columns

- .fitted  Predicted (or fitted) value
- .lower  Lower bound on interval
- .upper  Upper bound on interval
- .se.fit  Standard error of the predicted (or fitted) value

See Also

tidy.spmodel() glance.spmodel() predict.spmodel()

Examples

```r
spmod <- splm(z ~ water + tarp,
  data = caribou,
  spcov_type = "exponential", xcoord = x, ycoord = y)
augment(spmod)
spmod_sulf <- splm(sulfate ~ 1, data = sulfate, spcov_type = "exponential")
augment(spmod_sulf)
augment(spmod_sulf, newdata = sulfate_preds)
# missingness in original data
spmod_seal <- spautor(log_trend ~ 1, data = seal, spcov_type = "car")
augment(spmod_seal)
augment(spmod_seal, newdata = spmod_seal$newdata)
```
## Description

A caribou forage experiment.

## Usage

```r
caribou
```

## Format

A `tibble` with 30 rows and 5 columns:

- **water**: A factor representing whether water was added. Takes values `N` (no water added) and `Y` (water added).
- **tarp**: A factor representing tarp cover. Takes values `clear` (a clear tarp), `shade` (a shade tarp), and `none` (no tarp).
- **z**: The percentage of nitrogen.
- **x**: The x-coordinate.
- **y**: The y-coordinate.

## Source

These data were provided by Elizabeth Lenart of the Alaska Department of Fish and Game. The data were used in the publication listed in References.

## References

Usage

## S3 method for class 'splm'
coef(object, type = "fixed", ...)  

## S3 method for class 'splm'
coefficients(object, type = "fixed", ...)  

## S3 method for class 'spautor'
coef(object, type = "fixed", ...)  

## S3 method for class 'spautor'
coefficients(object, type = "fixed", ...)  

## S3 method for class 'spglm'
coef(object, type = "fixed", ...)  

## S3 method for class 'spglm'
coefficients(object, type = "fixed", ...)  

## S3 method for class 'spgautor'
coef(object, type = "fixed", ...)  

## S3 method for class 'spgautor'
coefficients(object, type = "fixed", ...)  

Arguments

object A fitted model object from splm(), spautor(), spglm(), or spgautor().  
type "fixed" for fixed effect coefficients, "spcov" for spatial covariance parameter coefficients, or "randcov" for random effect variance coefficients. Defaults to "fixed". If type = "spcov", the coefficient vector is an spcov_params() object (which means that has class matching the spatial covariance function used).  
... Other arguments. Not used (needed for generic consistency).

Value

A named vector of coefficients.

Examples

spmod <- splm(z ~ water + tarp,  
data = caribou,  
spcov_type = "exponential", xcoord = x, ycoord = y  
)  
coef(spmod)  
coefficients(spmod)  
coef(spmod, type = "spcov")
Confidence intervals for fitted model parameters

Description
Computes confidence intervals for one or more parameters in a fitted model object.

Usage
```r
## S3 method for class 'splm'
confint(object, parm, level = 0.95, ...)

## S3 method for class 'spautor'
confint(object, parm, level = 0.95, ...)

## S3 method for class 'spglm'
confint(object, parm, level = 0.95, ...)

## S3 method for class 'spgautor'
confint(object, parm, level = 0.95, ...)
```

Arguments
- `object`: A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`.
- `parm`: A specification of which parameters are to be given confidence intervals (a character vector of names). If missing, all parameters are considered.
- `level`: The confidence level required. The default is 0.95.
- `...`: Other arguments. Not used (needed for generic consistency).

Value
Gaussian-based confidence intervals (two-sided and equal-tailed) for the fixed effect coefficients based on the confidence level specified by `level`. For `spglm()` or `spgautor()` fitted model objects, confidence intervals are on the link scale.

Examples
```r
spmod <- splm(z ~ water + tarp,
  data = caribou,
  spcov_type = "exponential", xcoord = x, ycoord = y
)
confint(spmod)
confint(spmod, parm = "waterY", level = 0.90)
```
Description

Compute the Cook's distance for each observation from a fitted model object.

Usage

```r
## S3 method for class 'splm'
cooks.distance(model, ...)  
## S3 method for class 'spautor'
cooks.distance(model, ...)  
## S3 method for class 'spglm'
cooks.distance(model, ...)  
## S3 method for class 'spgautor'
cooks.distance(model, ...)  
```

Arguments

- `model` A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`.
- `...` Other arguments. Not used (needed for generic consistency).

Details

Cook's distance measures the influence of an observation on a fitted model object. If an observation is influential, its omission from the data noticeably impacts parameter estimates. The larger the Cook's distance, the larger the influence.

Value

A vector of Cook's distance values for each observation from the fitted model object.

See Also

- `augment.spmodel()`, `hatvalues.spmodel()`, `influence.spmodel()`, `residuals.spmodel()`

Examples

```r
spmod <- splm(z ~ water + tarp,  
data = caribou,  
spcov_type = "exponential", xcoord = x, ycoord = y)
cooks.distance(spmod)
```
Description

Create a covariance matrix from a fitted model object.

Usage

covmatrix(object, newdata, ...)

## S3 method for class 'splm'
covmatrix(object, newdata, ...)

## S3 method for class 'spautor'
covmatrix(object, newdata, ...)

## S3 method for class 'spglm'
covmatrix(object, newdata, ...)

## S3 method for class 'spgautor'
covmatrix(object, newdata, ...)

Arguments

- **object**: A fitted model object (e.g., `splm()`, `spautor()`, `spglm()`, or `spgautor()`).
- **newdata**: If omitted, the covariance matrix of the observed data is returned. If provided, newdata is a data frame or sf object that contains coordinate information required to construct the covariance between newdata and the observed data. If a data frame, newdata must contain variables that represent coordinates having the same name as the coordinates from the observed data used to fit object. If an sf object, coordinates are obtained from the geometry of newdata.
- **...**: Other arguments. Not used (needed for generic consistency).

Value

If newdata is omitted, the covariance matrix of the observed data, which has dimension n x n, where n is the sample size used to fit object. If newdata is provided, the covariance matrix between the unobserved (new) data and the observed data, which has dimension m x n, where m is the number of new observations and n is the sample size used to fit object.

Examples

```r
spmod <- splm(z ~ water + tarp,
              data = caribou,
              spcov_type = "exponential", xcoord = x, ycoord = y
             )
covmatrix(spmod)
```
## Description

Returns the deviance of a fitted model object.

## Usage

```r
## S3 method for class 'splm'
deviance(object, ...)

## S3 method for class 'spautor'
deviance(object, ...)

## S3 method for class 'spglm'
deviance(object, ...)

## S3 method for class 'spgautor'
deviance(object, ...)
```

## Arguments

- **object**: A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`, where `estmethod` is "ml" or "reml".
- **...**: Other arguments. Not used (needed for generic consistency).

## Details

For objects estimated using "ml" or "reml", the deviance is twice the difference in log-likelihoods between the saturated (perfect-fit) model and the fitted model.

## Value

The deviance.

## Examples

```r
spmod <- splm(z ~ water + tarp,
              data = caribou,
              spcov_type = "exponential", xcoord = x, ycoord = y
             )
deviance(spmod)
```
dispersion_initial

Create a dispersion parameter initial object

Description

Create a dispersion parameter initial object that specifies initial and/or known values to use while estimating the dispersion parameter with spglm() or spgautor().

Usage

dispersion_initial(family, dispersion, known)

Arguments

- **family**: The generalized linear model family describing the distribution of the response variable to be used. "poisson", "nbinomial", "binomial", "beta", "Gamma", and "inverse.gaussian".
- **dispersion**: The value of the dispersion parameter.
- **known**: A character vector indicating whether the dispersion parameter is to be assumed known. The value "dispersion" or "given" is assumes the dispersion parameter is known.

Details

The dispersion_initial list is later passed to spglm() or spgautor().

The variance function of an individual $y$ (given $\mu$) for each generalized linear model family is given below:

- family: $\text{Var}(y)$
  - poisson: $\mu \phi$
  - nbinomial: $\mu + \mu^2 / \phi$
  - binomial: $n\mu(1 - \mu)\phi$
  - beta: $\mu(1 - \mu)/(1 + \phi)$
  - Gamma: $\mu^2 / \phi$
  - inverse.gaussian: $\mu^2 / \phi$

The parameter $\phi$ is a dispersion parameter that influences $\text{Var}(y)$. For the poisson and binomial families, $\phi$ is always one. Note that this inverse Gaussian parameterization is different than a standard inverse Gaussian parameterization, which has variance $\mu^3 / \lambda$. Setting $\phi = \lambda / \mu$ yields our parameterization, which is preferred for computational stability. Also note that the dispersion parameter is often defined in the literature as $V(\mu)\phi$, where $V(\mu)$ is the variance function of the mean. We do not use this parameterization, which is important to recognize while interpreting dispersion parameter estimates using spglm() or spgautor(). For more on generalized linear model constructions, see McCullagh and Nelder (1989).
dispersion_params

Value
A list with two elements: initial and is_known. initial is a named numeric vector indicating the dispersion parameters with a specified initial and/or known value. is_known is a named numeric vector indicating whether the dispersion parameters in initial is known or not. The class of the list matches the value given to the family argument.

References

Examples
# known dispersion value 1
dispersion_initial("nbinomial", dispersion = 1, known = "dispersion")

---
dispersion_params Create a dispersion parameter object

Description
Create a dispersion parameter object for use with other functions.

Usage
dispersion_params(family, dispersion)

Arguments
family The generalized linear model family describing the distribution of the response variable to be used. "poisson", "nbinomial", "binomial", "beta", "Gamma", and "inverse.gaussian".
dispersion The value of the dispersion parameter.

Details
The variance function of an individual $y$ (given $\mu$) for each generalized linear model family is given below:

- family: $\text{Var}(y)$
- poisson: $\mu\phi$
- nbinomial: $\mu + \mu^2/\phi$
- binomial: $n\mu(1 - \mu)\phi$
- beta: $\mu(1 - \mu)/(1 + \phi)$
- Gamma: $\mu^2/\phi$
- inverse.gaussian: $\mu^2/\phi$
The parameter $\phi$ is a dispersion parameter that influences $\text{Var}(y)$. For the poisson and binomial families, $\phi$ is always one. Note that this inverse Gaussian parameterization is different than a standard inverse Gaussian parameterization, which has variance $\mu^3/\lambda$. Setting $\phi = \lambda/\mu$ yields our parameterization, which is preferred for computational stability. Also note that the dispersion parameter is often defined in the literature as $V(\mu)/\phi$, where $V(\mu)$ is the variance function of the mean. We do not use this parameterization, which is important to recognize while interpreting dispersion parameter estimates using `spglm()` or `spgautor()`. For more on generalized linear model constructions, see McCullagh and Nelder (1989).

Value

A named numeric vector with class `family` containing the dispersion.

References


Examples

dispersion_params("beta", dispersion = 1)

Description

Compute the empirical semivariogram for varying bin sizes and cutoff values.

Usage

esv(
  formula, 
  data,
  xcoord,
  ycoord,
  dist_matrix,
  bins = 15,
  cutoff,
  partition_factor
)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>formula</code></td>
<td>A formula describing the fixed effect structure.</td>
</tr>
<tr>
<td><code>data</code></td>
<td>A data frame or <code>sf</code> object containing the variables in <code>formula</code> and geographic information.</td>
</tr>
<tr>
<td><code>xcoord</code></td>
<td>Name of the variable in <code>data</code> representing the x-coordinate. Can be quoted or unquoted. Not required if <code>data</code> is an <code>sf</code> object.</td>
</tr>
</tbody>
</table>
ycoord: Name of the variable in data representing the y-coordinate. Can be quoted or unquoted. Not required if data is an sf object.

dist_matrix: A distance matrix to be used instead of providing coordinate names.

bins: The number of equally spaced bins. The default is 15.

cutoff: The maximum distance considered. The default is half the diagonal of the bounding box from the coordinates.

partition_factor: An optional formula specifying the partition factor. If specified, semivariances are only computed for observations sharing the same level of the partition factor.

Details

The empirical semivariogram is a tool used to visualize and model spatial dependence by estimating the semivariance of a process at varying distances. For a constant-mean process, the semivariance at distance \( h \) is denoted \( \gamma(h) \) and defined as \( 0.5 \times \text{Var}(z_1 - z_2) \). Under second-order stationarity, \( \gamma(h) = \text{Cov}(0) - \text{Cov}(h) \), where \( \text{Cov}(h) \) is the covariance function at distance \( h \). Typically the residuals from an ordinary least squares fit defined by formula are second-order stationary with mean zero. These residuals are used to compute the empirical semivariogram. At a distance \( h \), the empirical semivariance is \( 1/N(h) \sum (r_1 - r_2)^2 \), where \( N(h) \) is the number of (unique) pairs in the set of observations whose distance separation is \( h \) and \( r_1 \) and \( r_2 \) are residuals corresponding to observations whose distance separation is \( h \). In spmodel, these distance bins actually contain observations whose distance separation is \( h \pm c \), where \( c \) is a constant determined implicitly by bins. Typically, only observations whose distance separation is below some cutoff are used to compute the empirical semivariogram (this cutoff is determined by cutoff).

When using splm() with estmethod as "sv-wls", the empirical semivariogram is calculated internally and used to estimate spatial covariance parameters.

Value

A data frame with distance bins (bins), the average distance (dist), the semivariance (gamma), and the number of (unique) pairs (np).

Examples

esv(sulfate ~ 1, sulfate)
Usage

## S3 method for class 'splm'
fitted(object, type = "response", ...)

## S3 method for class 'splm'
fitted.values(object, type = "response", ...)

## S3 method for class 'spautor'
fitted(object, type = "response", ...)

## S3 method for class 'spautor'
fitted.values(object, type = "response", ...)

## S3 method for class 'spglm'
fitted(object, type = "response", ...)

## S3 method for class 'spglm'
fitted.values(object, type = "response", ...)

## S3 method for class 'spgautor'
fitted(object, type = "response", ...)

## S3 method for class 'spgautor'
fitted.values(object, type = "response", ...)

Arguments

object A fitted model object from splm(), spautor(), spglm(), or spgautor().
type "response" for fitted values of the response, "spcov" for fitted values of the spatial random errors, or "randcov" for fitted values of the random effects. If from spglm() or spgautor(), "link" for fitted values on the link scale. The default is "response".
... Other arguments. Not used (needed for generic consistency).

Details

When type is "response", the fitted values for each observation are the standard fitted values \( \hat{X}_i \hat{\beta} \). When type is "spcov" the fitted values for each observation are (generally) the best linear unbiased predictors of the spatial dependent and spatial independent random error. When type is "randcov", the fitted values for each level of each random effect are (generally) the best linear unbiased predictors of the corresponding random effect. The fitted values for type "spcov" and "randcov" can generally be used to check assumptions for each component of the fitted model object (e.g., check a Gaussian assumption).

Value

The fitted values according to type.
Examples

```r
spmod <- splm(z ~ water + tarp,
data = caribou,
  spcov_type = "exponential", xcoord = x, ycoord = y
)
fitted(spmod)
fitted.values(spmod)
fitted(spmod, type = "spcov")
```

---

**Description**

Return formula used by a fitted model object.

**Usage**

```r
## S3 method for class 'spml'
formula(x, ...)

## S3 method for class 'spautor'
formula(x, ...)

## S3 method for class 'spglm'
formula(x, ...)

## S3 method for class 'spgautore'
formula(x, ...)
```

**Arguments**

- `x` A fitted model object from `spml()`, `spautor()`, `spglm()`, or `spgautore()`.
- `...` Other arguments. Not used (needed for generic consistency).

**Value**

The formula used by a fitted model object.

**Examples**

```r
spmod <- splm(z ~ water + tarp,
data = caribou,
  spcov_type = "exponential", xcoord = x, ycoord = y
)
formula(spmod)
```
Glance at a fitted model object

Description

Returns a row of model summaries from a fitted model object. Glance returns the same number of columns for all models and estimation methods. If a particular summary is undefined for a model or estimation method (e.g., likelihood statistics for estimation methods "sv-wls" or "sv-cl" of splm() objects), NA is returned for that summary.

Usage

## S3 method for class 'splm'
glance(x, ...)

## S3 method for class 'spautor'
glance(x, ...)

## S3 method for class 'spglm'
glance(x, ...)

## S3 method for class 'spgautor'
glance(x, ...)

Arguments

x

A fitted model object from splm(), spautor(), spglm(), or spgautor().

... Other arguments. Not used (needed for generic consistency).

Value

A single-row tibble with columns

- n The sample size.
- p The number of fixed effects.
- npar The number of estimated covariance parameters.
- value The optimized value of the fitting function
- AIC The AIC.
- AICc The AICc.
- logLik The log-likelihood
- deviance The deviance.
- pseudo.r.squared The pseudo r-squared
### glances

**Glance at many fitted model objects**

#### Description

`glances()` repeatedly calls `glance()` on several fitted model objects and binds the output together, sorted by a column of interest.

#### Usage

```r
glances(object, ..., sort_by = "AICc", decreasing = FALSE)
```

```r
## S3 method for class 'splm'
glances(object, ..., sort_by = "AICc", decreasing = FALSE)
```

```r
## S3 method for class 'spautor'
glances(object, ..., sort_by = "AICc", decreasing = FALSE)
```

```r
## S3 method for class 'splm_list'
glances(object, ..., sort_by = "AICc", decreasing = FALSE)
```

```r
## S3 method for class 'spautor_list'
glances(object, ..., sort_by = "AICc", decreasing = FALSE)
```

```r
## S3 method for class 'spglm'
glances(object, ..., sort_by = "AICc", decreasing = FALSE)
```

```r
## S3 method for class 'spgautor'
glances(object, ..., sort_by = "AICc", decreasing = FALSE)
```

```r
## S3 method for class 'spglm_list'
glances(object, ..., sort_by = "AICc", decreasing = FALSE)
```

```r
## S3 method for class 'spgautor_list'
glances(object, ..., sort_by = "AICc", decreasing = FALSE)
```

#### Examples

```r
spmod <- splm(z ~ water + tarp, 
  data = caribou,
  spcov_type = "exponential", xcoord = x, ycoord = y
)
glance(spmod)
```

```r
```
hatvalues.spmodel

Arguments

object A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`.

... Additional fitted model objects. Ignored if object has class `splm_list`, `spautor_list`, `spglm_list`, or `spgautor_list`.

sort_by Sort by a glance statistic (i.e., the name of a column output from `glance()` or the order of model input (`sort_by = "order"`). The default is "AICc".

decreasing Should `sort_by` be decreasing or not? The default is `FALSE`.

Value

A tibble where each row represents the output of `glance()` for each fitted model object.

Examples

```r
lmod <- splm(z ~ water + tarp,
             data = caribou,
             spcov_type = "none"
)
smod <- splm(z ~ water + tarp,
             data = caribou,
             spcov_type = "exponential", xcoord = x, ycoord = y
)
glances(lmod, smod)
glances(lmod, smod, sort_by = "logLik", decreasing = TRUE)
```

hatvalues.spmodel Compute leverage (hat) values

Description

Compute the leverage (hat) value for each observation from a fitted model object.

Usage

```r
## S3 method for class 'splm'
hatvalues(model, ...)

## S3 method for class 'spautor'
hatvalues(model, ...)

## S3 method for class 'spglm'
hatvalues(model, ...)

## S3 method for class 'spgautor'
hatvalues(model, ...)
```
Arguments

model A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`.

... Other arguments. Not used (needed for generic consistency).

Details

Leverage values measure how far an observation’s explanatory variables are relative to the average of the explanatory variables. In other words, observations with high leverage are typically considered to have an extreme or unusual combination of explanatory variables. Leverage values are the diagonal of the hat (projection) matrix. The larger the hat value, the larger the leverage.

Value

A vector of leverage (hat) values for each observation from the fitted model object.

See Also

`augment.spmodel()` `cooks.distance.spmodel()` `influence.spmodel()` `residuals.spmodel()`

Examples

```r
spmod <- splm(z ~ water + tarp,
  data = caribou,
  spcov_type = "exponential", xcoord = x, ycoord = y
)
hatvalues(spmod)
```

---

influence.spmodel Regression diagnostics

Description

Provides basic quantities which are used in forming a wide variety of diagnostics for checking the quality of fitted model objects.

Usage

```r
## S3 method for class 'splm'
influence(model, ...)

## S3 method for class 'spautor'
influence(model, ...)

## S3 method for class 'spglm'
influence(model, ...)

## S3 method for class 'spgautor'
influence(model, ...)
```
Arguments

model  A fitted model object from \texttt{splm()}, \texttt{spautor()}, \texttt{spglm()}, or \texttt{spgautor()}.

...  Other arguments. Not used (needed for generic consistency).

Details

This function calls \texttt{residuals.spmodel()}, \texttt{hatvalues.spmodel()}, and \texttt{cooks.distance.spmodel()} and puts the results into a tibble. It is primarily used when calling \texttt{augment.spmodel()}.

Value

A tibble with residuals (\texttt{.resid}), leverage values (\texttt{.hat}), cook's distance (\texttt{.cooksd}), and standardized residuals (\texttt{.std.resid}).

See Also

\texttt{augment.spmodel()}, \texttt{cooks.distance.spmodel()}, \texttt{hatvalues.spmodel()}, \texttt{residuals.spmodel()}

Examples

```r
spmod <- splm(z ~ water + tarp,
              data = caribou,
              spcov_type = "exponential", xcoord = x, ycoord = y
)
influence(spmod)
```

---

labels.spmodel  \textit{Find labels from object}

Description

Find a suitable set of labels from a fitted model object.

Usage

```r
## S3 method for class 'splm'
labels(object, ...)

## S3 method for class 'spautor'
labels(object, ...)

## S3 method for class 'spglm'
labels(object, ...)

## S3 method for class 'spgautor'
labels(object, ...)
```
Arguments

object      A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`.  
...          Other arguments. Not used (needed for generic consistency).

Value

A character vector containing the terms used for the fixed effects from a fitted model object.

Examples

```r
spmod <- splm(z ~ water + tarp,  
data = caribou,  
spcov_type = "exponential", xcoord = x, ycoord = y  
)
labels(spmod)
```

Description

Find the log-likelihood of a fitted model when `estmethod` is "ml" or "reml".

Usage

```r
## S3 method for class 'splm'
logLik(object, ...)

## S3 method for class 'spautor'
logLik(object, ...)

## S3 method for class 'spglm'
logLik(object, ...)

## S3 method for class 'spgautor'
logLik(object, ...)
```

Arguments

object      A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()` where  
estmethod is "ml" or "reml".  
...          Other arguments. Not used (needed for generic consistency).

Value

The log-likelihood.
Examples

```r
spmod <- splm(z ~ water + tarp,
             data = caribou,
             spcov_type = "exponential", xcoord = x, ycoord = y
          )
logLik(spmod)
```

Perform leave-one-out cross validation

Description

Perform leave-one-out cross validation with options for computationally efficient approximations for big data.

Usage

```r
loocv(object, ...)
```

## S3 method for class 'splm'
```r
loocv(object, cv_predict = FALSE, se.fit = FALSE, local, ...)
```

## S3 method for class 'spautor'
```r
loocv(object, cv_predict = FALSE, se.fit = FALSE, local, ...)
```

## S3 method for class 'spglm'
```r
loocv(object, cv_predict = FALSE, se.fit = FALSE, local, ...)
```

## S3 method for class 'spgautor'
```r
loocv(object, cv_predict = FALSE, se.fit = FALSE, local, ...)
```

Arguments

- `object`: A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`.
- `...`: Other arguments. Not used (needed for generic consistency).
- `cv_predict`: A logical indicating whether the leave-one-out fitted values should be returned. Defaults to FALSE. If object is from `spglm()` or `spgautor()`, the fitted values returned are on the link scale.
- `se.fit`: A logical indicating whether the leave-one-out prediction standard errors should be returned. Defaults to FALSE. If object is from `spglm()` or `spgautor()`, the standard errors correspond to the fitted values returned on the link scale.
- `local`: A list or logical. If a list, specific list elements described in `predict.spmodel()` control the big data approximation behavior. If a logical, TRUE chooses default list elements for the list version of local as specified in `predict.spmodel()`. Defaults to FALSE, which performs exact computations.
Details

Each observation is held-out from the data set and the remaining data are used to make a prediction for the held-out observation. This is compared to the true value of the observation and several fit statistics are computed: bias, mean-squared-prediction error (MSPE), root-mean-squared-prediction error (RMSPE), and the squared correlation (cor2) between the observed data and leave-one-out predictions (regarded as a prediction version of r-squared appropriate for comparing across spatial and nonspatial models). Generally, bias should be near zero for well-fitting models. The lower the MSPE and RMSPE, the better the model fit (according to the leave-out-out criterion). The higher the cor2, the better the model fit (according to the leave-out-out criterion). cor2 is not returned when object was fit using spglm() or spgautor(), as it is only applicable here for linear models.

Value

If cv.predict = FALSE and se.fit = FALSE, a fit statistics tibble (with bias, MSPE, RMSPE, and cor2; see Details). If cv.predict = TRUE or se.fit = TRUE, a list with elements: stats, a fit statistics tibble (with bias, MSPE, RMSPE, and cor2; see Details); cv.predict, a numeric vector with leave-one-out predictions for each observation (if cv.predict = TRUE); and se.fit, a numeric vector with leave-one-out prediction standard errors for each observation (if se.fit = TRUE).

Examples

```r
spmod <- splm(z ~ water + tarp,
               data = caribou,
               spcov_type = "exponential", xcoord = x, ycoord = y
) 
loocv(spmod)
loocv(spmod, cv_predict = TRUE, se.fit = TRUE)
```

---

**model.frame.spmodel**

Extract the model frame from a fitted model object

**Description**

Extract the model frame from a fitted model object.

**Usage**

```r
## S3 method for class 'splm'
model.frame(formula, ...)

## S3 method for class 'spautor'
model.frame(formula, ...)

## S3 method for class 'spglm'
model.frame(formula, ...)

## S3 method for class 'spgautor'
model.frame(formula, ...)
```
model.matrix.spmodel

Arguments

  formula  A fitted model object from `splm()`, `spautor()`, `spgautor()`, or `spgautor()`.
  ...      Other arguments. Not used (needed for generic consistency).

Value

  A model frame that contains the variables used by the formula for the fitted model object.

See Also

  `stats::model.frame()`

Examples

```r
spmod <- splm(z ~ water + tarp,
  data = caribou,
  spcov_type = "exponential", xcoord = x, ycoord = y
)
model.frame(spmod)
```

model.matrix.spmodel  Extract the model matrix from a fitted model object

Description

  Extract the model matrix (X) from a fitted model object.

Usage

```r
## S3 method for class 'splm'
model.matrix(object, ...)

## S3 method for class 'spautor'
model.matrix(object, ...)

## S3 method for class 'spgautor'
model.matrix(object, ...)

## S3 method for class 'spglm'
model.matrix(object, ...)
```

Arguments

  object  A fitted model object from `splm()`, `spautor()`, `spgautor()`, or `spgautor()`.
  ...    Other arguments. Not used (needed for generic consistency).
Value

The model matrix (of the fixed effects), whose rows represent observations and whose columns represent explanatory variables corresponding to each fixed effect.

See Also

stats::model.matrix()

Examples

spmod <- splm(z ~ water + tarp,
  data = caribou,
  spcov_type = "exponential", xcoord = x, ycoord = y
)
model.matrix(spmod)

---

**moose**

* Moose counts and presence in Alaska, USA

Description

Moose counts and presence in Alaska, USA.

Usage

moose

Format

An sf object with 218 rows and 5 columns.

- elev: The elevation.
- strat: A factor representing strata (used for sampling). Can take values L and M.
- count: The count (number) of moose observed.
- presence: A binary factor representing whether no moose were observed (value 0) or at least one moose was observed (value 1).
- geometry: POINT geometry representing coordinates in an Alaska Albers projection (EPSG: 3338).

Source

Alaska Department of Fish and Game, Division of Wildlife Conservation has released this data set under the CC0 license.
**moose_preds**

*Locations at which to predict moose counts and presence in Alaska, USA*

**Description**

Locations at which to predict moose counts and presence in Alaska, USA.

**Usage**

`moose_preds`

**Format**

An `sf` object with 100 rows and 3 columns.

- `elev`: The elevation.
- `strat`: A factor representing strata (used for sampling). Can take values L and M.
- `geometry`: `POINT` geometry representing coordinates in an Alaska Albers projection (EPSG: 3338).

**Source**

Alaska Department of Fish and Game, Division of Wildlife Conservation has released this data set under the CC0 license.

---

**moss**

*Heavy metals in mosses near a mining road in Alaska, USA*

**Description**

Heavy metals in mosses near a mining road in Alaska, USA.

**Usage**

`moss`

**Format**

An `sf` object with 365 rows and 10 columns:

- `sample`: A factor with a sample identifier. Some samples were replicated in the field or laboratory. As a result, there are 318 unique sample identifiers.
- `field_dup`: A factor representing field duplicate. Takes values 1 and 2.
- `lab_rep`: A factor representing laboratory replicate. Takes values 1 and 2.
- sideroad: A factor representing direction relative to the haul road. Takes values N (north of the haul road) and S (south of the haul road).
- log_dist2road: The log of distance (in meters) to the haul road.
- log_Zn: The log of zinc concentration in moss tissue (mg/kg).
- geometry: POINT geometry representing coordinates in an Alaska Albers projection (EPSG: 3338).

Source

Data were obtained from Peter Neitlich and Linda Hasselbach of the National Park Service. Data were used in the publications listed in References.

References


plot.spmodel

Plot fitted model diagnostics

Description

Plot fitted model diagnostics such as residuals vs fitted values, quantile-quantile, scale-location, Cook’s distance, residuals vs leverage, Cook’s distance vs leverage, a fitted spatial covariance function, and a fitted anisotropic level curve of equal correlation.

Usage

```r
## S3 method for class 'splm'
plot(x, which, ...)

## S3 method for class 'spautor'
plot(x, which, ...)

## S3 method for class 'spglm'
plot(x, which, ...)

## S3 method for class 'spgautor'
plot(x, which, ...)
```
Arguments

- **x**: A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`.

- **which**: An integer vector taking on values between 1 and 7, which indicates the plots to return. Available plots are described in Details. If `which` has length greater than one, additional plots are stepped through in order using <Return>. The default for `splm()` and `spglm()` fitted model objects is `which = c(1, 2, 7)`. The default for `spautor()` and `spgautor()` fitted model objects is `which = c(1, 2)`.

- **...**: Other arguments passed to other methods.

Details

For all fitted model objects, the values of `which` make the corresponding plot:

- 1: Standardized residuals vs fitted values (of the response)
- 2: Normal quantile-quantile plot of standardized residuals
- 3: Scale-location plot of standardized residuals
- 4: Cook’s distance
- 5: Standardized residuals vs leverage
- 6: Cook’s distance vs leverage

For `splm()` and `spglm()` fitted model objects, there are two additional values of `which`:

- 7: Fitted spatial covariance function vs distance
- 8: Fitted anisotropic level curve of equal correlation

Value

No return value. Function called for plotting side effects.

Examples

```r
spmod <- splm(z ~ water + tarp,
               data = caribou,
               spcov_type = "exponential", xcoord = x, ycoord = y
            )
plot(spmod)
plot(spmod, which = c(1, 2, 4, 6))
```
**predict.spmodel**

Model predictions (Kriging)

**Description**

Predicted values and intervals based on a fitted model object.

**Usage**

```r
## S3 method for class 'splm'
predict(
  object,
  newdata,
  se.fit = FALSE,
  interval = c(\"none\", \"confidence\", \"prediction\"),
  level = 0.95,
  local,
  ...
)

## S3 method for class 'spautor'
predict(
  object,
  newdata,
  se.fit = FALSE,
  interval = c(\"none\", \"confidence\", \"prediction\"),
  level = 0.95,
  local,
  ...
)

## S3 method for class 'splm_list'
predict(
  object,
  newdata,
  se.fit = FALSE,
  interval = c(\"none\", \"confidence\", \"prediction\"),
  level = 0.95,
  local,
  ...
)

## S3 method for class 'spautor_list'
predict(
  object,
  newdata,
  se.fit = FALSE,
```
predict.spmodel

interval = c("none", "confidence", "prediction"),
level = 0.95,
local,
...
)

## S3 method for class 'splmRF'
predict(object, newdata, local, ...)

## S3 method for class 'spautorRF'
predict(object, newdata, local, ...)

## S3 method for class 'splmRF_list'
predict(object, newdata, local, ...)

## S3 method for class 'spautorRF_list'
predict(object, newdata, local, ...)

## S3 method for class 'spglm'
predict(
  object,
  newdata,
  type = c("link", "response"),
  se.fit = FALSE,
  interval = c("none", "confidence", "prediction"),
  newdata_size,
  level = 0.95,
  local,
  var_correct = TRUE,
  ...
)

## S3 method for class 'spgautor'
predict(
  object,
  newdata,
  type = c("link", "response"),
  se.fit = FALSE,
  interval = c("none", "confidence", "prediction"),
  newdata_size,
  level = 0.95,
  local,
  var_correct = TRUE,
  ...
)

## S3 method for class 'spglm_list'
predict(
Arguments

- **object**: A fitted model object.
- **newdata**: A data frame or sf object in which to look for variables with which to predict. If a data frame, newdata must contain all variables used by `formula(object)` and all variables representing coordinates. If an sf object, newdata must contain all variables used by `formula(object)` and coordinates are obtained from the geometry of newdata. If omitted, missing data from the fitted model object are used.
- **se.fit**: A logical indicating if standard errors are returned. The default is FALSE.
- **interval**: Type of interval calculation. The default is "none". Other options are "confidence" (for confidence intervals) and "prediction" (for prediction intervals).
- **level**: Tolerance/confidence level. The default is 0.95.
- **local**: A optional logical or list controlling the big data approximation. If omitted, local is set to TRUE or FALSE based on the sample size of the fitted model object and/or the prediction size of newdata – if the sample size or prediction size exceeds 5000, local is set to TRUE, otherwise it is set to FALSE. If FALSE, no big data approximation is implemented. If a list is provided, the following arguments detail the big data approximation:
  - **method**: The big data approximation method. If method = "all", all observations are used and size is ignored. If method = "distance", the
size data observations closest (in terms of Euclidean distance) to the observation requiring prediction are used. If method = "covariance", the size data observations with the highest covariance with the observation requiring prediction are used. If random effects and partition factors are not used in estimation and the spatial covariance function is monotone decreasing, "distance" and "covariance" are equivalent. The default is "covariance". Only used with models fit using splm() or spglm().

• size: The number of data observations to use when method = "distance" or "covariance". The default is 100. Only used with models fit using splm() or spglm().
• parallel: If TRUE, parallel processing via the parallel package is automatically used. The default is FALSE.
• ncores: If parallel = TRUE, the number of cores to parallelize over. The default is the number of available cores on your machine.

When local is a list, at least one list element must be provided to initialize default arguments for the other list elements. If local is TRUE, defaults for local are chosen such that local is transformed into list(size = 100, method = "covariance", parallel = FALSE).

... Other arguments. Only used for models fit using splmRF() or spautorRF() where ... indicates other arguments to ranger::predict.ranger().

type The scale (response or link) of predictions obtained using spglm() or spgautor objects.

newdata_size The size value for each observation in newdata used when predicting for the binomial family.

var_correct A logical indicating whether to return the corrected prediction variances when predicting via models fit using spglm() or spgautor(). The default is TRUE.

Details

For splm and spautor objects, the (empirical) best linear unbiased predictions (i.e., Kriging predictions) at each site are returned when interval is "none" or "prediction" alongside standard errors. Prediction intervals are also returned if interval is "prediction". When interval is "confidence", the estimated mean is returned alongside standard errors and confidence intervals for the mean. For splm_list and spautor_list objects, predictions and associated intervals and standard errors are returned for each list element.

For splmRF or spautorRF objects, random forest spatial residual model predictions are computed by combining the random forest prediction with the (empirical) best linear unbiased prediction for the residual. Fox et al. (2020) call this approach random forest regression Kriging. For splmRF_list or spautorRF objects, predictions are returned for each list element.

Value

For splm or spautor objects, if se.fit is FALSE, predict() returns a vector of predictions or a matrix of predictions with column names fit, lwr, and upr if interval is "confidence" or "prediction". If se.fit is TRUE, a list with the following components is returned:

• fit: vector or matrix as above
• se.fit: standard error of each fit

For splm_list or spautor_list objects, a list that contains relevant quantities for each list element.

For splmRF or spautorRF objects, a vector of predictions. For splmRF_list or spautorRF_list objects, a list that contains relevant quantities for each list element.

References


Examples

```r
spmod <- splm(sulfate ~ 1,
               data = sulfate,
               spcov_type = "exponential", xcoord = x, ycoord = y)
predict(spmod, sulfate_preds)
predict(spmod, sulfate_preds, interval = "prediction")
augment(spmod, newdata = sulfate_preds, interval = "prediction")

sulfate$var <- rnorm(NROW(sulfate)) # add noise variable
sulfate_preds$var <- rnorm(NROW(sulfate_preds)) # add noise variable
sprfmod <- splmRF(sulfate ~ var, data = sulfate, spcov_type = "exponential")
predict(sprfmod, sulfate_preds)
```

print.spmodel

*Print values*

Description

Print fitted model objects and summaries.

Usage

```r
## S3 method for class 'splm'
print(x, digits = max(3L,getOption("digits") - 3L), ...)

## S3 method for class 'spautor'
print(x, digits = max(3L,getOption("digits") - 3L), ...)

## S3 method for class 'summary.splm'
print(x,
      digits = max(3L,getOption("digits") - 3L),
      signif.stars = getOption("show.signif.stars"),
```
## S3 method for class 'summary.spautor'
print(
  x,
  digits = max(3L, getOption("digits") - 3L),
  signif.stars = getOption("show.signif.stars"),
  ...
)

## S3 method for class 'anova.splm'
print(
  x,
  digits = max(getOption("digits") - 2L, 3L),
  signif.stars = getOption("show.signif.stars"),
  ...
)

## S3 method for class 'anova.spautor'
print(
  x,
  digits = max(getOption("digits") - 2L, 3L),
  signif.stars = getOption("show.signif.stars"),
  ...
)

## S3 method for class 'spglm'
print(x, digits = max(3L, getOption("digits") - 3L), ...)

## S3 method for class 'spgautor'
print(x, digits = max(3L, getOption("digits") - 3L), ...)

## S3 method for class 'summary.spglm'
print(
  x,
  digits = max(3L, getOption("digits") - 3L),
  signif.stars = getOption("show.signif.stars"),
  ...
)

## S3 method for class 'summary.spgautor'
print(
  x,
  digits = max(3L, getOption("digits") - 3L),
  signif.stars = getOption("show.signif.stars"),
  ...
## S3 method for class 'anova.spglm'
print(
    x,
    digits = max(getOption("digits") - 2L, 3L),
    signif.stars = getOption("show.signif.stars"),
    ...
)

## S3 method for class 'anova.spgautor'
print(
    x,
    digits = max(getOption("digits") - 2L, 3L),
    signif.stars = getOption("show.signif.stars"),
    ...
)

### Arguments

- **x**
  - A fitted model object from `splm()`, `spgautor()`, `spglm()`, or `spgautor()` or output from `summary(x)` or `anova(x)`.
- **digits**
  - The number of significant digits to use when printing.
- **...**
  - Other arguments passed to or from other methods.
- **signif.stars**
  - Logical. If TRUE, significance stars are printed for each coefficient.

### Value

Printed fitted model objects and summaries with formatting.

### Examples

```r
spmod <- splm(z ~ water + tarp,
    data = caribou,
    spcov_type = "exponential", xcoord = x, ycoord = y
)
print(spmod)
print(summary(spmod))
print(anova(spmod))
```

---

### Description

Compute a pseudo r-squared for a fitted model object.
Usage

pseudoR2(object, ...)

## S3 method for class 'splm'
pseudoR2(object, adjust = FALSE, ...)

## S3 method for class 'spautor'
pseudoR2(object, adjust = FALSE, ...)

## S3 method for class 'spglm'
pseudoR2(object, adjust = FALSE, ...)

## S3 method for class 'spgautor'
pseudoR2(object, adjust = FALSE, ...)

Arguments

object A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`.
...

Other arguments. Not used (needed for generic consistency).

adjust A logical indicating whether the pseudo r-squared should be adjusted to account for the number of explanatory variables. The default is `FALSE`.

Details

Several pseudo r-squared statistics exist for in the literature. We define this pseudo r-squared as one minus the ratio of the deviance of a full model relative to the deviance of a null (intercept only) model. This pseudo r-squared can be viewed as a generalization of the classical r-squared definition seen as one minus the ratio of error sums of squares from the full model relative to the error sums of squares from the null model. If adjusted, the adjustment is analogous to the the classical r-squared adjustment.

Value

The pseudo r-squared as a numeric vector.

Examples

spmod <- splm(z ~ water + tarp,
             data = caribou,
             spcov_type = "exponential", xcoord = x, ycoord = y
)
pseudoR2(spmod)
**randcov_initial**

Create a random effects covariance parameter initial object

**Description**
Create a random effects (co)variance parameter initial object that specifies initial and/or known values to use while estimating random effect variances with modeling functions.

**Usage**
randcov_initial(..., known)

**Arguments**
- ... Arguments to randcov_params()
- known A character vector indicating which random effect variances are to be assumed known. The value "given" is shorthand for assuming all random effect variances given to randcov_initial() are assumed known.

**Details**
A random effect is specified as $Zu$, where $Z$ is the random effects design matrix and $u$ is the random effect. The covariance of $Zu$ is $\sigma^2 ZZ^T$, where $\sigma^2$ is the random effect variance, and $Z^T$ is the transpose of $Z$.

**Value**
A list with two elements: `initial` and `is_known`. `initial` is a named numeric vector indicating the random effect variances with specified initial and/or known values. `is_known` is a named logical vector indicating whether the random effect variances in `initial` are known or not.

**Examples**
- randcov_initial(group = 1)
- randcov_initial(group = 1, known = "group")

---

**randcov_params**

Create a random effects covariance parameter object

**Description**
Create a random effects covariance parameter object for use with other functions.

**Usage**
randcov_params(..., nn)

**Description**
Create a random effects covariance parameter object for use with other functions.
Arguments

... A named vector (or vectors) whose names represent the name of each random effect and whose values represent the variance of each random effect. If unnamed, nm is used to set names.

nm A character vector of names to assign to ....

Details

Names of the random effects should match eligible names given to random in modeling functions. While with the random argument to these functions, an intercept is implicitly assumed, with randcov_params, an intercept must be explicitly specified. That is, while with random, x | group is shorthand for (1 | group) + (x | group), with randcov_params, x | group implies just x | group, which means that if 1 | group is also desired, it must be explicitly specified.

Value

A named numeric vector of random effect covariance parameters.

Examples

randcov_params(group = 1, subgroup = 2)
randcov_params(1, 2, nm = c("group", "subgroup"))
# same as
randcov_params("1 | group" = 1, "1 | subgroup" = 2)

residuals.spmodel Extract fitted model residuals

Description

Extract residuals from a fitted model object. resid is an alias.

Usage

## S3 method for class 'splm'
residuals(object, type = "response", ...)

## S3 method for class 'splm'
resid(object, type = "response", ...)

## S3 method for class 'spautor'
rstandard(model, ...)

## S3 method for class 'spautor'
residuals(object, type = "response", ...)
### Arguments

- **object**: A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`.
- **type**: "response" for response residuals, "pearson" for Pearson residuals, or "standardized" for standardized residuals. For `splm()` and `spautor()` fitted model objects, the default is "response". For `spglm()` and `spgautor()` fitted model objects, deviance residuals are also available ("deviance") and are the default residual type.
- **...**: Other arguments. Not used (needed for generic consistency).
- **model**: A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`.

### Details

The response residuals are taken as the response minus the fitted values for the response: \( y - X \hat{\beta} \). The Pearson residuals are the response residuals pre-multiplied by their inverse square root. The standardized residuals are Pearson residuals divided by the square root of one minus the leverage (hat) value. The standardized residuals are often used to check model assumptions, as they have mean zero and variance approximately one.

`rstandard()` is an alias for `residuals(model, type = "standardized")`.

### Value

The residuals as a numeric vector.
See Also

augment.spmodel() cooks.distance.spmodel() hatvalues.spmodel() influence.spmodel()

Examples

```r
spmod <- splm(z ~ water + tarp,
    data = caribou,
    spcov_type = "exponential", xcoord = x, ycoord = y
)
residuals(spmod)
resid(spmod)
residuals(spmod, type = "pearson")
residuals(spmod, type = "standardized")
rstandard(spmod)
```

---

**seal**

*Estimated harbor-seal trends from abundance data in southeast Alaska, USA*

**Description**

Estimated harbor-seal trends from abundance data in southeast Alaska, USA.

**Usage**

`seal`

**Format**

A `sf` object with 62 rows and 2 columns:

- `log_trend`: The log of the estimated harbor-seal trends from abundance data.
- `geometry`: POLYGON geometry representing polygons in an Alaska Albers projection (EPSG: 3338).

**Source**

These data were collected by the Polar Ecosystem Program of the Marine Mammal Laboratory of the Alaska Fisheries Science Center of NOAA Fisheries. The data were used in the publication listed in References.

**References**

spautor

Fit spatial autoregressive models

Description
Fit spatial linear models for areal data (i.e., spatial autoregressive models) using a variety of estimation methods, allowing for random effects, partition factors, and row standardization.

Usage
spautor(
  formula,
  data,
  spcov_type,
  spcov_initial,
  estmethod = "reml",
  random,
  randcov_initial,
  partition_factor,
  W,
  row_st = TRUE,
  M,
  range_positive = TRUE,
  ...
)

Arguments

formula A two-sided linear formula describing the fixed effect structure of the model, with the response to the left of the \( \sim \) operator and the terms, separated by + operators, on the right.

data A data frame or sf object that contains the variables in fixed, random, and partition_factor, as well as potentially geographical information.

spcov_type The spatial covariance type. Available options include "car" and "sar". Parameterizations of each spatial covariance type are available in Details. When spcov_type is specified, relevant spatial covariance parameters are assumed unknown, requiring estimation. spcov_type is not required (and is ignored) if spcov_initial is provided. Multiple values can be provided in a character vector. Then spautor() is called iteratively for each element and a list is returned for each model fit. The default for spcov_type is "car".

spcov_initial An object from spcov_initial() specifying initial and/or known values for the spatial covariance parameters. Not required if spcov_type is provided. Multiple spcov_initial() objects can be provided in a list. Then spautor() is called iteratively for each element and a list is returned for each model fit.

estmethod The estimation method. Available options include "reml" for restricted maximum likelihood and "ml" for maximum likelihood. The default is "reml".
random

A one-sided linear formula describing the random effect structure of the model. Terms are specified to the right of the ~ operator. Each term has the structure $x_1 + \ldots + x_n \mid g_1/\ldots/g_m$, where $x_1 + \ldots + x_n$ specifies the model for the random effects and $g_1/\ldots/g_m$ is the grouping structure. Separate terms are separated by + and must generally be wrapped in parentheses. Random intercepts are added to each model implicitly when at least one other variable is defined. If a random intercept is not desired, this must be explicitly defined (e.g., $x_1 + \ldots + x_n - 1 \mid g_1/\ldots/g_m$). If only a random intercept is desired for a grouping structure, the random intercept must be specified as $1 \mid g_1/\ldots/g_m$. Note that $g_1/\ldots/g_m$ is shorthand for $(1 \mid g_1/\ldots/g_m)$. If only random intercepts are desired and the shorthand notation is used, parentheses can be omitted.

randcov_initial

An optional object specifying initial and/or known values for the random effect variances.

partition_factor

A one-sided linear formula with a single term specifying the partition factor. The partition factor assumes observations from different levels of the partition factor are uncorrelated.

$W$

Weight matrix specifying the neighboring structure used. Not required if data is an sf polygon object, as $W$ is calculated internally using queen contiguity. If calculated internally, $W$ is computed using sf::st_intersects().

row_st

A logical indicating whether row standardization be performed on $W$. The default is TRUE.

$M$

$M$ matrix satisfying the car symmetry condition. The car symmetry condition states that $(I - \text{range} \ast W)^{-1}M$ is symmetric, where $I$ is an identity matrix, range is a constant that controls the spatial dependence, $W$ is the weights matrix, and $^{-1}$ represents the inverse operator. $M$ is required for car models when $W$ is provided and row_st is FALSE. When $M$, is required, the default is the identity matrix. $M$ must be diagonal or given as a vector or one-column matrix assumed to be the diagonal.

range_positive

Whether the range should be constrained to be positive. The default is TRUE.

... Other arguments to stats::optim().

Details

The spatial linear model for areal data (i.e., spatial autoregressive model) can be written as $y = X\beta + \tau + \epsilon$, where $X$ is the fixed effects design matrix, $\beta$ are the fixed effects, $\tau$ is random error that is spatially dependent, and $\epsilon$ is random error that is spatially independent. Together, $\tau$ and $\epsilon$ are modeled using a spatial covariance function, expressed as $de \ast R + ie \ast I$, where $de$ is the dependent error variance, $R$ is a matrix that controls the spatial dependence structure among observations, $ie$ is the independent error variance, and $I$ is an identity matrix. Note that $de$ and $ie$ must be non-negative while range must be between the reciprocal of the maximum eigenvalue of $W$ and the reciprocal of the minimum eigenvalue of $W$.

spcov_type Details: Parametric forms for $R$ are given below:

- car: $(I - \text{range} \ast W)^{-1}M$, weights matrix $W$, symmetry condition matrix $M$
- sar: $[(I - \text{range} \ast W)(I - \text{range} \ast W)^T]^{-1}$, weights matrix $W$, $^T$ indicates matrix transpose
If there are observations with no neighbors, they are given a unique variance parameter called extra, which must be non-negative.

**estmethod** Details: The various estimation methods are

- **reml**: Maximize the restricted log-likelihood.
- **ml**: Maximize the log-likelihood.

By default, all spatial covariance parameters except ie as well as all random effect variance parameters are assumed unknown, requiring estimation. ie is assumed zero and known by default (in contrast to models fit using splm(), where ie is assumed unknown by default). To change this default behavior, specify spcov_initial (an NA value for ie in spcov_initial to assume ie is unknown, requiring estimation).

**random** Details: If random effects are used, the model can be written as \( y = X\beta + Z_1u_1 + \ldots Z_ju_j + \tau + \epsilon \), where each \( Z \) is a random effects design matrix and each \( u \) is a random effect.

**partition_factor** Details: The partition factor can be represented in matrix form as \( P \), where elements of \( P \) equal one for observations in the same level of the partition factor and zero otherwise. The covariance matrix involving only the spatial and random effects components is then multiplied element-wise (Hadamard product) by \( P \), yielding the final covariance matrix.

Observations with NA response values are removed for model fitting, but their values can be predicted afterwards by running predict(object). This is the only way to perform prediction for spautor() models (i.e., the prediction locations must be known prior to estimation).

**Value**

A list with many elements that store information about the fitted model object. If spcov_type or spcov_initial are length one, the list has class spautor. Many generic functions that summarize model fit are available for spautor objects, including AIC, AICc, anova, augment, coef, cooks.distance, covmatrix, deviance, fitted, formula, glance, glances, hatvalues, influence, labels, logLik, loocv, model.frame, model.matrix, plot, predict, print, pseudoR2, summary, terms, tidy, update, varcomp, and vcov. If spcov_type or spcov_initial are length greater than one, the list has class spautor_list and each element in the list has class spautor. glances can be used to summarize spautor_list objects, and the aforementioned spautor generics can be used on each individual list element (model fit).

**Note**

This function does not perform any internal scaling. If optimization is not stable due to large extremely large variances, scale relevant variables so they have variance 1 before optimization.

**Examples**

```r
spmod <- spautor(log_trend ~ 1, data = seal, spcov_type = "car")
summary(spmod)
```
spautorRF | *Fit random forest spatial residual models*

**Description**

Fit random forest residual spatial linear models for areal data (i.e., spatial autoregressive models) using random forest to fit the mean and a spatial linear model to fit the residuals. The spatial linear model fit to the residuals can incorporate a variety of estimation methods, allowing for random effects, partition factors, and row standardization.

**Usage**

```r
spautorRF(formula, data, ...)
```

**Arguments**

- `formula`: A two-sided linear formula describing the fixed effect structure of the model, with the response to the left of the `~` operator and the terms on the right, separated by `+` operators.
- `data`: A data frame or `sf` object object that contains the variables in `fixed`, `random`, and `partition_factor` as well as geographical information. If an `sf` object is provided with `POINT` geometries, the x-coordinates and y-coordinates are used directly. If an `sf` object is provided with `POLYGON` geometries, the x-coordinates and y-coordinates are taken as the centroids of each polygon.
- `...`: Additional named arguments to `ranger::ranger()` or `spautor()`.

**Details**

The random forest residual spatial linear model is described by Fox et al. (2020). A random forest model is fit to the mean portion of the model specified by `formula` using `ranger::ranger()`. Residuals are computed and used as the response variable in an intercept-only spatial linear model fit using `spautor()`. This model object is intended for use with `predict()` to perform prediction, also called random forest regression Kriging.

**Value**

A list with several elements to be used with `predict()`. These elements include the function call (named `call`), the random forest object fit to the mean (named `ranger`), the spatial linear model object fit to the residuals (named `spautor` or `spautor_list`), and an object can contain data for locations at which to predict (called `newdata`). The `newdata` object contains the set of observations in `data` whose response variable is `NA`. If `spcov_type` or `spcov_initial` (which are passed to `spautor()`) are length one, the list has class `spautorRF` and the spatial linear model object fit to the residuals is called `spautor`, which has class `spautor`. If `spcov_type` or `spcov_initial` are length greater than one, the list has class `spautorRF_list` and the spatial linear model object fit to the residuals is called `spautor_list`, which has class `spautor_list` and contains several objects, each with class `spautor`. 
References

Examples

```r
seal$var <- rnorm(NROW(seal)) # add noise variable
sprfmod <- spautorRF(log_trend ~ var, data = seal, spcov_type = "car")
predict(sprfmod)
```

**spcov_initial**

Create a spatial covariance parameter initial object

**Description**
Create a spatial covariance parameter initial object that specifies initial and/or known values to use while estimating spatial covariance parameters with `splm()`, `spglm()`, `spautor()`, or `spgautor()`.

**Usage**

```r
spcov_initial(spcov_type, de, ie, range, extra, rotate, scale, known)
```

**Arguments**

- **spcov_type**

- **de**
  The spatially dependent (correlated) random error variance. Commonly referred to as a partial sill.

- **ie**
  The spatially independent (uncorrelated) random error variance. Commonly referred to as a nugget.

- **range**
  The correlation parameter.

- **extra**
  An extra covariance parameter used when spcov_type is "matern", "cauchy", "pexponential", "car", or "sar".

- **rotate**
  Anisotropy rotation parameter (from 0 to π radians). Not used if spcov_type is "car" or "sar".

- **scale**
  Anisotropy scale parameter (from 0 to 1). Not used if spcov_type is "car" or "sar".

- **known**
  A character vector indicating which spatial covariance parameters are to be assumed known. The value "given" is shorthand for assuming all spatial covariance parameters given to spcov_initial() are assumed known.
Details

The spcov_initial list is later passed to splm(), spgml(), spautor(), or spgautor(). NA values can be given for ie, rotate, and scale, which lets these functions find initial values for parameters that are sometimes otherwise assumed known (e.g., rotate and scale with splm() and spgml() and ie with spautor() and spgautor()). The spatial covariance functions can be generally expressed as $de * R + ie * I$, where $de$ is $de$ above, $R$ is a matrix that controls the spatial dependence structure among observations, $h$, $ie$ is $ie$ above, and $I$ is an identity matrix. Note that $de$ and $ie$ must be non-negative while range must be positive, except when spcov_type is car or sar, in which case range must be between the reciprocal of the maximum eigenvalue of $W$ and the reciprocal of the minimum eigenvalue of $W$. Parametric forms for $R$ are given below, where $\eta = h/range$:

- exponential: $\exp(-\eta)$
- spherical: $(1 - 1.5\eta + 0.5\eta^2) * I(h <= range)$
- gaussian: $\exp(-\eta^2)$
- triangular: $(1 - \eta) * I(h <= range)$
- circular: $(1 - (2/\pi) * (m * \sqrt{(1 - m^2)} + \sin^{-1}(m))) * I(h <= range), m = \min(\eta, 1)$
- cubic: $(1 - 7\eta^2 + 8.75\eta^3 - 3.5\eta^5 + 0.75\eta^7) * I(h <= range)$
- pentaspherical: $(1 - 1.875\eta + 1.25\eta^3 - 0.375\eta^5) * I(h <= range)$
- cosine: $\cos(\eta)$
- wave: $\sin(\eta) / \eta * I(h > 0) + I(h = 0)$
- jbessel: $B_j(h * range)$, $B_j$ is Bessel-J function
- gravity: $(1 + \eta^2)^{-0.5}$
- rquad: $(1 + \eta^2)^{-1}$
- magnetic: $(1 + \eta^2)^{-1.5}$
- matern: $2^{1-extr\alpha} / \Gamma(extr\alpha) * \alpha^{extr\alpha} * Bk(\alpha, extr\alpha), \alpha = (2 extr\alpha * \eta)^{0.5}, Bk$ is Bessel-K function wit order $1/5 \leq extr \leq 5$
- cauchy: $(1 + \eta^2)^{-extr\alpha}, extr > 0$
- pexponential: $\exp(h^{extr\alpha}/range), 0 < extr \leq 2$
- car: $(I - range * W)^{-1} * M$, weights matrix $W$, symmetry condition matrix $M$, observations with no neighbors are given a unique variance parameter called $extr\alpha$, $extr \geq 0$.
- sar: $[(I - range * W)(I - range * W)^T]^{-1}$, weights matrix $W, T$ indicates matrix transpose, observations with no neighbors are given a unique variance parameter called $extr\alpha$, $extr \geq 0$.
- none: 0

All spatial covariance functions are valid in one spatial dimension. All spatial covariance functions except triangular and cosine are valid in two dimensions.

When the spatial covariance function is car or sar, $extr\alpha$ represents the variance parameter for the observations in $W$ without at least one neighbor (other than itself) – these are called unconnected observations. $extr\alpha$ is only used if there is at least one unconnected observation.
**spcov_params**

**Value**

A list with two elements: `initial` and `is_known`. `initial` is a named numeric vector indicating the spatial covariance parameters with specified initial and/or known values. `is_known` is a named numeric vector indicating whether the spatial covariance parameters in `initial` are known or not. The class of the list matches the value given to the `spcov_type` argument.

**Examples**

```r
# known de value 1 and initial range value 0.4
spcov_initial("exponential", de = 1, range = 0.4, known = c("de"))
# known ie value 0 and known range value 1
spcov_initial("gaussian", ie = 0, range = 1, known = c("given"))
# ie given NA
spcov_initial("car", ie = NA)
```

---

**spcov_params**

Create a spatial covariance parameter object

**Description**

Create a spatial covariance parameter object for use with other functions.

**Usage**

```r
spcov_params(spcov_type, de, ie, range, extra, rotate = 0, scale = 1)
```

**Arguments**

- **de**: The spatially dependent (correlated) random error variance. Commonly referred to as a partial sill.
- **ie**: The spatially independent (uncorrelated) random error variance. Commonly referred to as a nugget.
- **range**: The correlation parameter.
- **extra**: An extra covariance parameter used when `spcov_type` is "matern", "cauchy", "pexponential", "car", or "sar".
- **rotate**: Anisotropy rotation parameter (from 0 to π radians). A value of 0 (the default) implies no rotation. Not used if `spcov_type` is "car" or "sar".
- **scale**: Anisotropy scale parameter (from 0 to 1). A value of 1 (the default) implies no scaling. Not used if `spcov_type` is "car" or "sar".
Details

Generally, all arguments to `spcov_params` must be specified, though default arguments are often chosen based on `spcov_type`. When `spcov_type` is `car` or `sar`, `ie` is assumed to be 0 unless specified otherwise. For full parameterizations of all spatial covariance functions, see `spcov_initial()`.

Value

A named numeric vector of spatial covariance parameters with class `spcov_type`.

Examples

```r
spcov_params("exponential", de = 1, ie = 1, range = 1)
```

---

**spgautor**  
*Fit spatial generalized autoregressive models*

Description

Fit spatial generalized linear models for areal data (i.e., spatial generalized autoregressive models) using a variety of estimation methods, allowing for random effects, partition factors, and row standardization.

Usage

```r
spgautor(
  formula,
  family,
  data,
  spcov_type,
  spcov_initial,
  dispersion_initial,
  estmethod = "reml",
  random,
  randcov_initial,
  partition_factor,
  W,
  row_st = TRUE,
  M,
  range_positive = TRUE,
  ...
)
```

Arguments

- `formula` A two-sided linear formula describing the fixed effect structure of the model, with the response to the left of the `~` operator and the terms, separated by `+` operators, on the right.
family
The generalized linear model family describing the distribution of the response variable to be used. Available options "poisson", "nbinomial", "binomial", "beta", "Gamma", and "inverse.gaussian". Can be quoted or unquoted. Note that the family argument only takes a single value, rather than the list structure used by stats::glm. See Details for more.

data
A data frame or sf object that contains the variables in fixed, random, and partition_factor, as well as potentially geographical information.

spcov_type
The spatial covariance type. Available options include "car" and "sar". Parameterizations of each spatial covariance type are available in Details. When spcov_type is specified, relevant spatial covariance parameters are assumed unknown, requiring estimation. spcov_type is not required (and is ignored) if spcov_initial is provided. Multiple values can be provided in a character vector. Then spgautor() is called iteratively for each element and a list is returned for each model fit. The default for spcov_type is "car".

spcov_initial
An object from spcov_initial() specifying initial and/or known values for the spatial covariance parameters. Not required if spcov_type is provided. Multiple spcov_initial() objects can be provided in a list. Then spgautor() is called iteratively for each element and a list is returned for each model fit.

dispersion_initial
An object from dispersion_initial() specifying initial and/or known values for the dispersion parameter for the "nbinomial", "beta", "Gamma", and "inverse.gaussian" families. family is ignored if dispersion_initial is provided.

estmethod
The estimation method. Available options include "reml" for restricted maximum likelihood and "ml" for maximum likelihood. The default is "reml".

random
A one-sided linear formula describing the random effect structure of the model. Terms are specified to the right of the ~ operator. Each term has the structure x1 + ... + xn | g1/.../gm, where x1 + ... + xn specifies the model for the random effects and g1/.../gm is the grouping structure. Separate terms are separated by + and must generally be wrapped in parentheses. Random intercepts are added to each model implicitly when at least one other variable is defined. If a random intercept is not desired, this must be explicitly defined (e.g., x1 + ... + xn - 1 | g1/.../gm). If only a random intercept is desired for a grouping structure, the random intercept must be specified as 1 | g1/.../gm. Note that g1/.../gm is shorthand for (1 | g1/.../gm). If only random intercepts are desired and the shorthand notation is used, parentheses can be omitted.

randcov_initial
An optional object specifying initial and/or known values for the random effect variances.

partition_factor
A one-sided linear formula with a single term specifying the partition factor. The partition factor assumes observations from different levels of the partition factor are uncorrelated.

W
Weight matrix specifying the neighboring structure used. Not required if data is an sf polygon object, as W is calculated internally using queen contiguity. If calculated internally, W is computed using sf::st_intersects().
row_st

A logical indicating whether row standardization be performed on W. The default is TRUE.

M

M matrix satisfying the car symmetry condition. The car symmetry condition states that \((I - \text{range} \ast W)^{-1}M\) is symmetric, where I is an identity matrix, range is a constant that controls the spatial dependence, W is the weights matrix, and \(^{-1}\) represents the inverse operator. M is required for car models when W is provided and row_st is FALSE. When M, is required, the default is the identity matrix. M must be diagonal or given as a vector or one-column matrix assumed to be the diagonal.

range_positive

Whether the range should be constrained to be positive. The default is TRUE.

... Other arguments to stats::optim().

Details

The spatial generalized linear model for areal data (i.e., spatial generalized autoregressive model) can be written as \(g(\mu) = \eta = X\beta + \tau + \epsilon\), where \(\mu\) is the expectation of the response \((y)\) given the random errors, \(g(.)\) is called a link function which links together the \(\mu\) and \(\eta\), \(X\) is the fixed effects design matrix, \(\beta\) are the fixed effects, \(\tau\) is random error that is spatially dependent, and \(\epsilon\) is random error that is spatially independent.

There are six generalized linear model families available: poisson assumes \(y\) is a Poisson random variable nbioomial assumes \(y\) is a negative binomial random variable, binomial assumes \(y\) is a binomial random variable, beta assumes \(y\) is a beta random variable, Gamma assumes \(y\) is a gamma random variable, and inverse.gaussian assumes \(y\) is an inverse Gaussian random variable.

The supports for \(y\) for each family are given below:

- family: support of \(y\)
- poisson: \(0 \leq y; y\) an integer
- nbinomial: \(0 \leq y; y\) an integer
- binomial: \(0 \leq y; y\) an integer
- beta: \(0 < y < 1\)
- Gamma: \(0 < y\)
- inverse.gaussian: \(0 < y\)

The generalized linear model families and the parameterizations of their link functions are given below:

- family: link function
- poisson: \(g(\mu) = log(\eta)\) (log link)
- nbinomial: \(g(\mu) = log(\eta)\) (log link)
- binomial: \(g(\mu) = log(\eta/(1 - \eta))\) (logit link)
- beta: \(g(\mu) = log(\eta/(1 - \eta))\) (logit link)
- Gamma: \(g(\mu) = log(\eta)\) (log link)
- inverse.gaussian: \(g(\mu) = log(\eta)\) (log link)
The variance function of an individual $y$ (given $\mu$) for each generalized linear model family is given below:

- family: $\text{Var}(y)$
- poisson: $\mu \phi$
- nbinomial: $\mu + \mu^2 / \phi$
- binomial: $n \mu (1 - \mu) \phi$
- beta: $\mu (1 - \mu) / (1 + \phi)$
- Gamma: $\mu^2 / \phi$
- inverse.gaussian: $\mu^2 / \phi$

The parameter $\phi$ is a dispersion parameter that influences $\text{Var}(y)$. For the poisson and binomial families, $\phi$ is always one. Note that this inverse Gaussian parameterization is different than a standard inverse Gaussian parameterization, which has variance $\mu^3 / \lambda$. Setting $\phi = \lambda / \mu$ yields our parameterization, which is preferred for computational stability. Also note that the dispersion parameter is often defined in the literature as $V(\mu) \phi$, where $V(\mu)$ is the variance function of the mean. We do not use this parameterization, which is important to recognize while interpreting dispersion parameter estimates. For more on generalized linear model constructions, see McCullagh and Nelder (1989).

Together, $\tau$ and $\epsilon$ are modeled using a spatial covariance function, expressed as $de \ast R + ie \ast I$, where $de$ is the dependent error variance, $R$ is a matrix that controls the spatial dependence structure among observations, $ie$ is the independent error variance, and $I$ is an identity matrix. Note that $de$ and $ie$ must be non-negative while range must be between the reciprocal of the maximum eigenvalue of $W$ and the reciprocal of the minimum eigenvalue of $W$. Recall that $\tau$ and $\epsilon$ are modeled on the link scale, not the inverse link (response) scale. Random effects are also modeled on the link scale.

**spcov_type** Details: Parametric forms for $R$ are given below:

- car: $(I - \text{range} \ast W)^{-1} M$, weights matrix $W$, symmetry condition matrix $M$
- sar: $[(I - \text{range} \ast W)(I - \text{range} \ast W)^T]^{-1}$, weights matrix $W$, $^T$ indicates matrix transpose

If there are observations with no neighbors, they are given a unique variance parameter called extra, which must be non-negative.

**estmethod** Details: The various estimation methods are

- reml: Maximize the restricted log-likelihood.
- ml: Maximize the log-likelihood.

Note that the likelihood being optimized is obtained using the Laplace approximation.

By default, all spatial covariance parameters except $ie$ as well as all random effect variance parameters are assumed unknown, requiring estimation. $ie$ is assumed zero and known by default (in contrast to models fit using `spglm()`, where $ie$ is assumed unknown by default). To change this default behavior, specify spcov_initial (an NA value for $ie$ in spcov_initial to assume $ie$ is unknown, requiring estimation).

**random** Details: If random effects are used, the model can be written as $y = X \beta + Z_1 u_1 + \ldots Z_j u_j + \tau + \epsilon$, where each $Z$ is a random effects design matrix and each $u$ is a random effect.
spglm

Details: The partition factor can be represented in matrix form as $P$, where elements of $P$ equal one for observations in the same level of the partition factor and zero otherwise. The covariance matrix involving only the spatial and random effects components is then multiplied element-wise (Hadamard product) by $P$, yielding the final covariance matrix.

Observations with NA response values are removed for model fitting, but their values can be predicted afterwards by running `predict(object)`. This is the only way to perform prediction for `spgautor()` models (i.e., the prediction locations must be known prior to estimation).

Value

A list with many elements that store information about the fitted model object. If `spcov_type` or `spcov_initial` are length one, the list has class `spgautor`. Many generic functions that summarize model fit are available for `spgautor` objects, including `AIC`, `AICc`, `anova`, `augment`, `coef`, `cooks.distance`, `covmatrix`, `deviance`, `fitted`, `formula`, `glance`, `glances`, `hatvalues`, `influence`, `labels`, `logLik`, `loocv`, `model.frame`, `model.matrix`, `plot`, `predict`, `print`, `pseudoR2`, `summary`, `terms`, `tidy`, `update`, `varcomp`, and `vcov`. If `spcov_type` or `spcov_initial` are length greater than one, the list has class `spgautor_list` and each element in the list has class `spgautor`. `glances` can be used to summarize `spgautor_list` objects, and the aforementioned `spgautor` generics can be used on each individual list element (model fit).

Note

This function does not perform any internal scaling. If optimization is not stable due to large extremely large variances, scale relevant variables so they have variance 1 before optimization.

References


Examples

```r
spgmod <- spgautor(I(log_trend^2) ~ 1, family = "Gamma", data = seal, spcov_type = "car")
summary(spgmod)
```

---

spglm

Fit spatial generalized linear models

Description

Fit spatial generalized linear models for point-referenced data (i.e., generalized geostatistical models) using a variety of estimation methods, allowing for random effects, anisotropy, partition factors, and big data methods.
Usage

```r
spglm(
  formula, 
  family, 
  data, 
  spcov_type, 
  xcoord, 
  ycoord, 
  spcov_initial, 
  dispersion_initial, 
  estmethod = "reml", 
  anisotropy = FALSE, 
  random, 
  randcov_initial, 
  partition_factor, 
  local, 
  ...
)
```

Arguments

- **formula**: A two-sided linear formula describing the fixed effect structure of the model, with the response to the left of the `~` operator and the terms on the right, separated by `+` operators.

- **family**: The generalized linear model family describing the distribution of the response variable to be used. Available options "poisson", "nbinomial", "binomial", "beta", "Gamma", and "inverse.gaussian". Can be quoted or unquoted. Note that the family argument only takes a single value, rather than the list structure used by `stats::glm`. See Details for more.

- **data**: A data frame or `sf` object object that contains the variables in `fixed`, `random`, and `partition_factor` as well as geographical information. If an `sf` object is provided with `POINT` geometries, the x-coordinates and y-coordinates are used directly. If an `sf` object is provided with `POLYGON` geometries, the x-coordinates and y-coordinates are taken as the centroids of each polygon.

- **spcov_type**: The spatial covariance type. Available options include "exponential", "spherical", "gaussian", "triangular", "circular", "cubic", "pentspherical", "cosine", "wave", "jbessel", "gravity", "rquad", "magnetic", "matern", "cauchy", "pexponential", and "none". Parameterizations of each spatial covariance type are available in Details. Multiple spatial covariance types can be provided as a character vector, and then `spglm()` is called iteratively for each element and a list is returned for each model fit. The default for `spcov_type` is "exponential". When `spcov_type` is specified, all unknown spatial covariance parameters are estimated. `spcov_type` is ignored if `spcov_initial` is provided.

- **xcoord**: The name of the column in `data` representing the x-coordinate. Can be quoted or unquoted. Not required if `data` is an `sf` object.
ycoord  The name of the column in data representing the y-coordinate. Can be quoted or unquoted. Not required if data is an sf object.

spcov_initial  An object from spcov_initial() specifying initial and/or known values for the spatial covariance parameters. Multiple spcov_initial() objects can be provided in a list. Then spglm() is called iteratively for each element and a list is returned for each model fit.

dispersion_initial  An object from dispersion_initial() specifying initial and/or known values for the dispersion parameter for the "nbinomial", "beta", "Gamma", and "inverse.gaussian" families. family is ignored if dispersion_initial is provided.

estmethod  The estimation method. Available options include "reml" for restricted maximum likelihood and "ml" for maximum likelihood. The default is "reml".

anisotropy  A logical indicating whether (geometric) anisotropy should be modeled. Not required if spcov_initial is provided with 1) rotate assumed unknown or assumed known and non-zero or 2) scale assumed unknown or assumed known and less than one. When anisotropy is TRUE, computational times can significantly increase. The default is FALSE.

random  A one-sided linear formula describing the random effect structure of the model. Terms are specified to the right of the ~ operator. Each term has the structure x1 + ... + xn | g1/.../gm, where x1 + ... + xn specifies the model for the random effects and g1/.../gm is the grouping structure. Separate terms are separated by + and must generally be wrapped in parentheses. Random intercepts are added to each model implicitly when at least one other variable is defined. If a random intercept is not desired, this must be explicitly defined (e.g., x1 + ... + xn - 1 | g1/.../gm). If only a random intercept is desired for a grouping structure, the random intercept must be specified as 1 | g1/.../gm. Note that g1/.../gm is shorthand for (1 | g1/.../gm). If only random intercepts are desired and the shorthand notation is used, parentheses can be omitted.

randcov_initial  An optional object specifying initial and/or known values for the random effect variances.

partition_factor  A one-sided linear formula with a single term specifying the partition factor. The partition factor assumes observations from different levels of the partition factor are uncorrelated.

local  An optional logical or list controlling the big data approximation. If omitted, local is set to TRUE or FALSE based on the sample size (the number of non-missing observations in data) – if the sample size exceeds 3,000, local is set to TRUE. Otherwise it is set to FALSE. If FALSE, no big data approximation is implemented. If a list is provided, the following arguments detail the big data approximation:

- index: The group indexes. Observations in different levels of index are assumed to be uncorrelated for the purposes of estimation. If index is not provided, it is determined by specifying method and either size or groups.
method: The big data approximation method used to determine index. Ignored if index is provided. If method = "random", observations are randomly assigned to index based on size. If method = "kmeans", observations assigned to index based on k-means clustering on the coordinates with groups clusters. The default is "kmeans". Note that both methods have a random component, which means that you may get different results from separate model fitting calls. To ensure consistent results, specify index or set a seed via base::set.seed().

- size: The number of observations in each index group when method is "random". If the number of observations is not divisible by size, some levels get size - 1 observations. The default is 100.
- groups: The number of index groups. If method is "random", size is ceiling(n/groups), where n is the sample size. Automatically determined if size is specified. If method is "kmeans", groups is the number of clusters.
- var_adjust: The approach for adjusting the variance-covariance matrix of the fixed effects. "none" for no adjustment, "theoretical" for the theoretically-correct adjustment, "pooled" for the pooled adjustment, and "empirical" for the empirical adjustment. The default is "theoretical".
- parallel: If TRUE, parallel processing via the parallel package is automatically used. The default is FALSE.
- ncores: If parallel = TRUE, the number of cores to parallelize over. The default is the number of available cores on your machine.

When local is a list, at least one list element must be provided to initialize default arguments for the other list elements. If local is TRUE, defaults for local are chosen such that local is transformed into list(size = 100, method = "kmeans", var_adjust = "theoretical", parallel = FALSE).

... Other arguments to esv() or stats::optim().

Details

The spatial generalized linear model for point-referenced data (i.e., generalized geostatistical model) can be written as \( g(\mu) = \eta = X\beta + \tau + \epsilon \), where \( \mu \) is the expectation of the response \( y \) given the random errors, \( g(.) \) is called a link function which links together the \( \mu \) and \( \eta \), \( X \) is the fixed effects design matrix, \( \beta \) are the fixed effects, \( \tau \) is random error that is spatially dependent, and \( \epsilon \) is random error that is spatially independent.

There are six generalized linear model families available: poisson assumes \( y \) is a Poisson random variable, nbinomial assumes \( y \) is a negative binomial random variable, binomial assumes \( y \) is a binomial random variable, beta assumes \( y \) is a beta random variable, Gamma assumes \( y \) is a gamma random variable, and inverse.gaussian assumes \( y \) is an inverse Gaussian random variable.

The supports for \( y \) for each family are given below:

- family: support of \( y \)
  - poisson: \( 0 \leq y; y \) an integer
  - nbinomial: \( 0 \leq y; y \) an integer
  - binomial: \( 0 \leq y; y \) an integer
• beta: $0 < y < 1$
• Gamma: $0 < y$
• inverse.gaussian: $0 < y$

The generalized linear model families and the parameterizations of their link functions are given below:

• family: link function
  • poisson: $g(\mu) = \log(\eta)$ (log link)
  • nbinomial: $g(\mu) = \log(\eta)$ (log link)
  • binomial: $g(\mu) = \log(\eta/(1 - \eta))$ (logit link)
  • beta: $g(\mu) = \log(\eta/(1 - \eta))$ (logit link)
  • Gamma: $g(\mu) = \log(\eta)$ (log link)
  • inverse.gaussian: $g(\mu) = \log(\eta)$ (log link)

The variance function of an individual $y$ (given $\mu$) for each generalized linear model family is given below:

• family: $\text{Var}(y)$
  • poisson: $\mu \phi$
  • nbinomial: $\mu + \mu^2/\phi$
  • binomial: $\eta \mu (1 - \mu) \phi$
  • beta: $\mu(1 - \mu)/(1 + \phi)$
  • Gamma: $\mu^2/\phi$
  • inverse.gaussian: $\mu^2/\phi$

The parameter $\phi$ is a dispersion parameter that influences $\text{Var}(y)$. For the poisson and binomial families, $\phi$ is always one. Note that this inverse Gaussian parameterization is different than a standard inverse Gaussian parameterization, which has variance $\mu^3/\lambda$. Setting $\phi = \lambda/\mu$ yields our parameterization, which is preferred for computational stability. Also note that the dispersion parameter is often defined in the literature as $V(\mu)\phi$, where $V(\mu)$ is the variance function of the mean. We do not use this parameterization, which is important to recognize while interpreting dispersion estimates. For more on generalized linear model constructions, see McCullagh and Nelder (1989).

Together, $\tau$ and $\epsilon$ are modeled using a spatial covariance function, expressed as $de*R + ie*I$, where $de$ is the dependent error variance, $R$ is a correlation matrix that controls the spatial dependence structure among observations, $ie$ is the independent error variance, and $I$ is an identity matrix. Recall that $\tau$ and $\epsilon$ are modeled on the link scale, not the inverse link (response) scale. Random effects are also modeled on the link scale.

spcov_type Details: Parametric forms for $R$ are given below, where $\eta = h/range$ for $h$ distance between observations:

• exponential: $exp(-\eta)$
• spherical: $(1 - 1.5\eta + 0.5\eta^3) * I(h <= range)$
• gaussian: $exp(-\eta^2)$
• triangular: $(1 - \eta) * I(h <= range)$
• circular: \((1 - (2/\pi) * (m * \sqrt{1 - m^2} + \sin^{-1}(m))) * I(h <= \text{range}), m = \min(\eta, 1)\)
• cubic: \((1 - 7\eta^2 + 8.75\eta^3 - 3.5\eta^5 + 0.75\eta^7) * I(h <= \text{range})\)
• pentaspherical: \((1 - 1.875\eta + 1.25\eta^3 - 0.375\eta^5) * I(h <= \text{range})\)
• cosine: \(\cos(\eta)\)
• wave: \(\sin(\eta) / \eta * I(h > 0) + I(h = 0)\)
• jbessel: \(Bj(h * \text{range}), Bj \text{ is Bessel-J function}\)
• gravity: \((1 + \eta^2)^{-0.5}\)
• rquad: \((1 + \eta^2)^{-1}\)
• magnetic: \((1 + \eta^2)^{-1.5}\)
• matern: \(2^{1 - \text{extra}} / \Gamma(\text{extra}) * \alpha^{\text{extra}} * Bk(\alpha, \text{extra}), \alpha = (2\text{extra} * \eta)^{0.5}, Bk \text{ is Bessel-K function with order 1/5 \leq \text{extra} \leq 5}\)
• cauchy: \((1 + \eta^2)^{-\text{extra}}, \text{extra} > 0\)
• pexponential: \(exp(h^{\text{extra}/\text{range}}), 0 < \text{extra} \leq 2\)
• none: 0

All spatial covariance functions are valid in one spatial dimension. All spatial covariance functions except triangular and cosine are valid in two dimensions.

**estmethod** Details: The various estimation methods are
- reml: Maximize the restricted log-likelihood.
- ml: Maximize the log-likelihood.

Note that the likelihood being optimized is obtained using the Laplace approximation.

**anisotropy** Details: By default, all spatial covariance parameters except rotate and scale as well as all random effect variance parameters are assumed unknown, requiring estimation. If either rotate or scale are given initial values other than 0 and 1 (respectively) or are assumed unknown in `spcov_initial()`, anisotropy is implicitly set to TRUE. (Geometric) Anisotropy is modeled by transforming a covariance function that decays differently in different directions to one that decays equally in all directions via rotation and scaling of the original coordinates. The rotation is controlled by the rotate parameter in \([0, \pi]\) radians. The scaling is controlled by the scale parameter in \([0, 1]\). The anisotropy correction involves first a rotation of the coordinates clockwise by rotate and then a scaling of the coordinates’ minor axis by the reciprocal of scale. The spatial covariance is then computed using these transformed coordinates.

**random** Details: If random effects are used, the model can be written as \(y = X\beta + Z1u1 + ...Zjuj + \tau + \epsilon\), where each Z is a random effects design matrix and each u is a random effect.

**partition_factor** Details: The partition factor can be represented in matrix form as \(P\), where elements of \(P\) equal one for observations in the same level of the partition factor and zero otherwise. The covariance matrix involving only the spatial and random effects components is then multiplied element-wise (Hadamard product) by \(P\), yielding the final covariance matrix.

**local** Details: The big data approximation works by sorting observations into different levels of an index variable. Observations in different levels of the index variable are assumed to be uncorrelated for the purposes of model fitting. Sparse matrix methods are then implemented for significant computational gains. Parallelization generally further speeds up computations when data sizes are larger than a few thousand. Both the "random" and "kmeans" values of method in local have
random components. That means you may get slightly different results when using the big data approximation and rerunning \texttt{spglm()} with the same code. For consistent results, either set a seed via \texttt{base::set.seed()} or specify \texttt{index} to \texttt{local}.

Observations with NA response values are removed for model fitting, but their values can be predicted afterwards by running \texttt{predict(object)}.

\subsection*{Value}

A list with many elements that store information about the fitted model object. If \texttt{spcov_type} or \texttt{spcov_initial} are length one, the list has class \texttt{spglm}. Many generic functions that summarize model fit are available for \texttt{spglm} objects, including \texttt{AIC}, \texttt{AICc}, \texttt{anova}, \texttt{coef}, \texttt{cooks.distance}, \texttt{covmatrix}, \texttt{deviance}, \texttt{fitted}, \texttt{formula}, \texttt{glance}, \texttt{glances}, \texttt{hatvalues}, \texttt{influence}, \texttt{labels}, \texttt{logLik}, \texttt{loocv}, \texttt{model.frame}, \texttt{model.matrix}, \texttt{plot}, \texttt{predict}, \texttt{print}, \texttt{pseudoR2}, \texttt{summary}, \texttt{terms}, \texttt{tidy}, \texttt{update}, \texttt{varcomp}, and \texttt{vcov}. If \texttt{spcov_type} or \texttt{spcov_initial} are length greater than one, the list has class \texttt{spglm_list} and each element in the list has class \texttt{spglm}. \texttt{glances} can be used to summarize \texttt{spglm_list} objects, and the aforementioned \texttt{spglm} generics can be used on each individual list element (model fit).

\subsection*{Note}

This function does not perform any internal scaling. If optimization is not stable due to large extremely large variances, scale relevant variables so they have variance 1 before optimization.

\subsection*{References}


\subsection*{Examples}

\begin{verbatim}
spgmod <- spglm(presence ~ elev,
               family = "binomial", data = moose,
               spcov_type = "exponential"
)
summary(spgmod)
\end{verbatim}

---

\textbf{splm}

\textit{Fit spatial linear models}

\subsection*{Description}

Fit spatial linear models for point-referenced data (i.e., geostatistical models) using a variety of estimation methods, allowing for random effects, anisotropy, partition factors, and big data methods.
Usage

```r
splm(
    formula,
    data,
    spcov_type,
    xcoord,
    ycoord,
    spcov_initial,
    estmethod = "reml",
    weights = "cressie",
    anisotropy = FALSE,
    random,
    randcov_initial,
    partition_factor,
    local,
    ...
)
```

Arguments

- `formula`: A two-sided linear formula describing the fixed effect structure of the model, with the response to the left of the `~` operator and the terms on the right, separated by `+` operators.

- `data`: A data frame or `sf` object object that contains the variables in `fixed`, `random`, and `partition_factor` as well as geographical information. If an `sf` object is provided with POINT geometries, the x-coordinates and y-coordinates are used directly. If an `sf` object is provided with POLYGON geometries, the x-coordinates and y-coordinates are taken as the centroids of each polygon.

- `spcov_type`: The spatial covariance type. Available options include "exponential", "spherical", "gaussian", "triangular", "circular", "cubic", "pentaspherical", "cosine", "wave", "jbessel", "gravity", "rquad", "magnetic", "matern", "cauchy", "pexponential", and "none". Parameterizations of each spatial covariance type are available in Details. Multiple spatial covariance types can be provided as a character vector, and then `splm()` is called iteratively for each element and a list is returned for each model fit. The default for `spcov_type` is "exponential". When `spcov_type` is specified, all unknown spatial covariance parameters are estimated. `spcov_type` is ignored if `spcov_initial` is provided.

- `xcoord`: The name of the column in `data` representing the x-coordinate. Can be quoted or unquoted. Not required if `data` is an `sf` object.

- `ycoord`: The name of the column in `data` representing the y-coordinate. Can be quoted or unquoted. Not required if `data` is an `sf` object.

- `spcov_initial`: An object from `spcov_initial()` specifying initial and/or known values for the spatial covariance parameters. Multiple `spcov_initial()` objects can be provided in a list. Then `splm()` is called iteratively for each element and a list is returned for each model fit.
estmethod: The estimation method. Available options include "reml" for restricted maximum likelihood, "ml" for maximum likelihood, "sv-wls" for semivariogram weighted least squares, and "sv-cl" for semivariogram composite likelihood. The default is "reml".

weights: Weights to use when estmethod is "sv-wls". Available options include "cressie", "cressie-dr", "cressie-nopairs", "cressie-dr-nopairs", "pairs", "pairs-invrd", "pairs-invrd", and "ols". Parameterizations for each weight are available in Details. The default is "cressie".

anisotropy: A logical indicating whether (geometric) anisotropy should be modeled. Not required if spcov_initial is provided with 1) rotate assumed unknown or assumed known and non-zero or 2) scale assumed unknown or assumed known and less than one. When anisotropy is TRUE, computational times can significantly increase. The default is FALSE.

random: A one-sided linear formula describing the random effect structure of the model. Terms are specified to the right of the ~ operator. Each term has the structure x1 + ... + xn | g1/.../gm, where x1 + ... + xn specifies the model for the random effects and g1/.../gm is the grouping structure. Separate terms are separated by + and must generally be wrapped in parentheses. Random intercepts are added to each model implicitly when at least one other variable is defined. If a random intercept is not desired, this must be explicitly defined (e.g., x1 + ... + xn - 1 | g1/.../gm). If only a random intercept is desired for a grouping structure, the random intercept must be specified as 1 | g1/.../gm. Note that g1/.../gm is shorthand for (1 | g1/.../gm). If only random intercepts are desired and the shorthand notation is used, parentheses can be omitted.

randcov_initial: An optional object specifying initial and/or known values for the random effect variances.

partition_factor: A one-sided linear formula with a single term specifying the partition factor. The partition factor assumes observations from different levels of the partition factor are uncorrelated.

local: An optional logical or list controlling the big data approximation. If omitted, local is set to TRUE or FALSE based on the sample size (the number of non-missing observations in data) – if the sample size exceeds 5,000, local is set to TRUE. Otherwise it is set to FALSE. local is also set to FALSE when spcov_type is "none" and there are no random effects specified via random. If FALSE, no big data approximation is implemented. If a list is provided, the following arguments detail the big data approximation:

- index: The group indexes. Observations in different levels of index are assumed to be uncorrelated for the purposes of estimation. If index is not provided, it is determined by specifying method and either size or groups.
- method: The big data approximation method used to determine index. Ignored if index is provided. If method = "random", observations are randomly assigned to index based on size. If method = "kmeans", observations assigned to index based on k-means clustering on the coordinates with groups clusters. The default is "kmeans". Note that both methods
have a random component, which means that you may get different results from separate model fitting calls. To ensure consistent results, specify index or set a seed via `base::set.seed()`.

- **size**: The number of observations in each index group when method is "random". If the number of observations is not divisible by size, some levels get `size - 1` observations. The default is 100.
- **groups**: The number of index groups. If method is "random", `size` is `ceiling(n/groups)`, where `n` is the sample size. Automatically determined if `size` is specified. If method is "kmeans", groups is the number of clusters.
- **var_adjust**: The approach for adjusting the variance-covariance matrix of the fixed effects. "none" for no adjustment, "theoretical" for the theoretically-correct adjustment, "pooled" for the pooled adjustment, and "empirical" for the empirical adjustment. The default is "theoretical".
- **parallel**: If TRUE, parallel processing via the parallel package is automatically used. The default is FALSE.
- **ncores**: If parallel = TRUE, the number of cores to parallelize over. The default is the number of available cores on your machine.

When `local` is a list, at least one list element must be provided to initialize default arguments for the other list elements. If `local` is TRUE, defaults for `local` are chosen such that `local` is transformed into `list(size = 100, method = "kmeans", var_adjust = "theoretical", parallel = FALSE).

Other arguments to `esv()` or `stats::optim()`.

### Details

The spatial linear model for point-referenced data (i.e., geostatistical model) can be written as $y = X\beta + \tau + \epsilon$, where $X$ is the fixed effects design matrix, $\beta$ are the fixed effects, $\tau$ is random error that is spatially dependent, and $\epsilon$ is random error that is spatially independent. Together, $\tau$ and $\epsilon$ are modeled using a spatial covariance function, expressed as $de \ast R + ie \ast I$, where $de$ is the dependent error variance, $R$ is a correlation matrix that controls the spatial dependence structure among observations, $ie$ is the independent error variance, and $I$ is an identity matrix.

**spcov_type** Details: Parametric forms for $R$ are given below, where $\eta = h/range$ for h distance between observations:

- **exponential**: $exp(-\eta)$
- **spherical**: $(1 - 1.5\eta + 0.5\eta^3) \ast I(h <= range)$
- **gaussian**: $exp(-\eta^2)$
- **triangular**: $(1 - \eta) \ast I(h <= range)$
- **circular**: $(1 - (2/\pi) \ast (m * sqrt(1 - m^2) + sin^{-1}(m))) \ast I(h <= range), m = min(\eta, 1)$
- **cubic**: $(1 - 7\eta^2 + 8.75\eta^3 - 3.5\eta^5 + 0.75\eta^7) \ast I(h <= range)$
- **pentaspherical**: $(1 - 1.875\eta + 1.25\eta^3 - 0.375\eta^5) \ast I(h <= range)$
- **cosine**: $cos(\eta)$
- **wave**: $sin(\eta)/\eta \ast I(h > 0) + I(h = 0)$
- **jbessel**: $B_j(h * range), B_j$ is Bessel-J function
• gravity: \((1 + \eta^2)^{-0.5}\)
• rquad: \((1 + \eta^2)^{-1}\)
• magnetic: \((1 + \eta^2)^{-1.5}\)
• matern: \(2^{1 - \text{extra}}/\Gamma(\text{extra}) \ast \alpha^{\text{extra}} \ast Bk(\alpha, \text{extra}), \alpha = (2\text{extra} \ast \eta)^{0.5}, Bk \text{ is Bessel-K function with order} 1/5 \leq \text{extra} \leq 5\)
• cauchy: \((1 + \eta^2)^{-\text{extra}}, \text{extra} > 0\)
• pexponential: \(\exp(h^{\text{extra}}/\text{range}), 0 < \text{extra} \leq 2\)
• none: 0

All spatial covariance functions are valid in one spatial dimension. All spatial covariance functions except triangular and cosine are valid in two dimensions.

**estmethod Details:** The various estimation methods are

- **reml:** Maximize the restricted log-likelihood.
- **ml:** Maximize the log-likelihood.
- **sv-wls:** Minimize the semivariogram weighted least squares loss.
- **sv-cl:** Minimize the semivariogram composite likelihood loss.

**anisotropy Details:** By default, all spatial covariance parameters except rotate and scale as well as all random effect variance parameters are assumed unknown, requiring estimation. If either rotate or scale are given initial values other than 0 and 1 (respectively) or are assumed unknown in `spcov_initial()`, anisotropy is implicitly set to TRUE. (Geometric) Anisotropy is modeled by transforming a covariance function that decays differently in different directions to one that decays equally in all directions via rotation and scaling of the original coordinates. The rotation is controlled by the rotate parameter in \([0, \pi]\) radians. The scaling is controlled by the scale parameter in \([0, 1]\). The anisotropy correction involves first a rotation of the coordinates clockwise by rotate and then a scaling of the coordinates’ minor axis by the reciprocal of scale. The spatial covariance is then computed using these transformed coordinates.

**random Details:** If random effects are used (the estimation method must be "reml" or "ml"), the model can be written as \(y = X \beta + Z_1 u_1 + \ldots Z_j u_j + \tau + \epsilon\), where each \(Z\) is a random effects design matrix and each \(u\) is a random effect.

**partition_factor Details:** The partition factor can be represented in matrix form as \(P\), where elements of \(P\) equal one for observations in the same level of the partition factor and zero otherwise. The covariance matrix involving only the spatial and random effects components is then multiplied element-wise (Hadamard product) by \(P\), yielding the final covariance matrix.

**local Details:** The big data approximation works by sorting observations into different levels of an index variable. Observations in different levels of the index variable are assumed to be uncorrelated for the purposes of model fitting. Sparse matrix methods are then implemented for significant computational gains. Parallelization generally further speeds up computations when data sizes are larger than a few thousand. Both the "random" and "kmeans" values of method in `local` have random components. That means you may get slightly different results when using the big data approximation and rerunning `splm()` with the same code. For consistent results, either set a seed via `base::set.seed()` or specify index to `local`.

Observations with NA response values are removed for model fitting, but their values can be predicted afterwards by running `predict(object)`. 
**Value**

A list with many elements that store information about the fitted model object. If spcov_type or spcov_initial are length one, the list has class `splm`. Many generic functions that summarize model fit are available for `splm` objects, including `AIC`, `AICc`, `anova`, `coef`, `cooks.distance`, `covmatrix`, `deviance`, `fitted`, `formula`, `glance`, `glances`, `hatvalues`, `influence`, `labels`, `logLik`, `loocv`, `model.frame`, `model.matrix`, `plot`, `predict`, `print`, `pseudoR2`, `summary`, `terms`, `tidy`, `update`, `varcomp`, and `vcov`. If spcov_type or spcov_initial are length greater than one, the list has class `splm_list` and each element in the list has class `splm`. `glances` can be used to summarize `splm_list` objects, and the aforementioned `splm` generics can be used on each individual list element (model fit).

**Note**

This function does not perform any internal scaling. If optimization is not stable due to large extremely large variances, scale relevant variables so they have variance 1 before optimization.

**Examples**

```r
spmod <- splm(z ~ water + tarp, data = caribou, 
              spcov_type = "exponential", xcoord = x, ycoord = y)
summary(spmod)
```

---

**splmRF**

*Fit random forest spatial residual models*

**Description**

Fit random forest spatial residual models for point-referenced data (i.e., geostatistical models) using random forest to fit the mean and a spatial linear model to fit the residuals. The spatial linear model fit to the residuals can incorporate variety of estimation methods, allowing for random effects, anisotropy, partition factors, and big data methods.

**Usage**

```r
splmRF(formula, data, ...)
```

**Arguments**

- `formula` A two-sided linear formula describing the fixed effect structure of the model, with the response to the left of the `~` operator and the terms on the right, separated by `+` operators.
- `data` A data frame or `sf` object object that contains the variables in `fixed`, `random`, and `partition_factor` as well as geographical information. If an `sf` object is provided with `POINT` geometries, the x-coordinates and y-coordinates are used directly. If an `sf` object is provided with `POLYGON` geometries, the x-coordinates and y-coordinates are taken as the centroids of each polygon.
- `...` Additional named arguments to `ranger::ranger()` or `splm()`.
Details

The random forest residual spatial linear model is described by Fox et al. (2020). A random forest model is fit to the mean portion of the model specified by formula using `ranger::ranger()`. Residuals are computed and used as the response variable in an intercept-only spatial linear model fit using `splm()`. This model object is intended for use with `predict()` to perform prediction, also called random forest regression Kriging.

Value

A list with several elements to be used with `predict()`. These elements include the function call (named `call`), the random forest object fit to the mean (named `ranger`), the spatial linear model object fit to the residuals (named `splm` or `splm_list`), and an object can contain data for locations at which to predict (called `newdata`). The `newdata` object contains the set of observations in `data` whose response variable is NA. If `spcov_type` or `spcov_initial` (which are passed to `splm()`) are length one, the list has class `splmRF` and the spatial linear model object fit to the residuals is called `splm`, which has class `splm`. If `spcov_type` or `spcov_initial` are length greater than one, the list has class `splmRF_list` and the spatial linear model object fit to the residuals is called `splm_list`, which has class `splm_list` and contains several objects, each with class `splm`.

An `splmRF` object to be used with `predict()`. There are three elements: ranger, the output from fitting the mean model with `ranger::ranger()`; splm, the output from fitting the spatial linear model to the ranger residuals; and `newdata`, the `newdata` object, if relevant.

Note

This function does not perform any internal scaling. If optimization is not stable due to large extremely large variances, scale relevant variables so they have variance 1 before optimization.

References


Examples

```r
sulfate$var <- rnorm(NROW(sulfate)) # add noise variable
sulfate_preds$var <- rnorm(NROW(sulfate_preds)) # add noise variable
sprfmod <- splmRF(sulfate ~ var, data = sulfate, spcov_type = "exponential")
predict(sprfmod, sulfate_preds)
```

```
---

<table>
<thead>
<tr>
<th>sprbeta</th>
<th>Simulate a spatial beta random variable</th>
</tr>
</thead>
</table>
```

Description

Simulate a spatial beta random variable with a specific mean and covariance structure.
Usage

```
sprbeta(
    spcov_params,
    dispersion = 1,
    mean = 0,
    samples = 1,
    data,
    randcov_params,
    partition_factor,
    ...
)
```

Arguments

- `spcov_params`: An `spcov_params()` object.
- `dispersion`: The dispersion value.
- `mean`: A numeric vector representing the mean. `mean` must have length 1 (in which case it is recycled) or length equal to the number of rows in `data`. The default is 0.
- `samples`: The number of independent samples to generate. The default is 1.
- `data`: A data frame or `sf` object containing spatial information.
- `randcov_params`: A `randcov_params()` object.
- `partition_factor`: A formula indicating the partition factor.
- `...`: Additional arguments passed to `sprnorm()`.

Details

The values of `spcov_params`, `mean`, and `randcov_params` are assumed to be on the link scale. They are used to simulate a latent normal (Gaussian) response variable using `sprnorm()`. This latent variable is the conditional mean used with `dispersion` to simulate a beta random variable.

Value

If `samples` is 1, a vector of random variables for each row of `data` is returned. If `samples` is greater than one, a matrix of random variables is returned, where the rows correspond to each row of `data` and the columns correspond to independent samples.

Examples

```
spcov_params_val <- spcov_params("exponential", de = 0.2, ie = 0.1, range = 1)
sprbeta(spcov_params_val, data = caribou, xcoord = x, ycoord = y)
sprbeta(spcov_params_val, samples = 5, data = caribou, xcoord = x, ycoord = y)
```
**sprbinom**

Simulate a spatial binomial random variable

**Description**

Simulate a spatial binomial random variable with a specific mean and covariance structure.

**Usage**

```r
sprbinom(
    spcov_params,
    mean = 0,
    size = 1,
    samples = 1,
    data,
    randcov_params,
    partition_factor,
    ...
)
```

**Arguments**

- `spcov_params`: An `spcov_params()` object.
- `mean`: A numeric vector representing the mean. `mean` must have length 1 (in which case it is recycled) or length equal to the number of rows in `data`. The default is 0.
- `size`: A numeric vector representing the sample size for each binomial trial. The default is 1, which corresponds to a Bernoulli trial for each observation.
- `samples`: The number of independent samples to generate. The default is 1.
- `data`: A data frame or `sf` object containing spatial information.
- `randcov_params`: A `randcov_params()` object.
- `partition_factor`: A formula indicating the partition factor.
- `...`: Additional arguments passed to `sprnorm()`.

**Details**

The values of `spcov_params`, `mean`, and `randcov_params` are assumed to be on the link scale. They are used to simulate a latent normal (Gaussian) response variable using `sprnorm()`. This latent variable is the conditional mean used with dispersion to simulate a binomial random variable.

**Value**

If `samples` is 1, a vector of random variables for each row of `data` is returned. If `samples` is greater than one, a matrix of random variables is returned, where the rows correspond to each row of `data` and the columns correspond to independent samples.
Examples

```r
spcov_params_val <- spcov_params("exponential", de = 0.2, ie = 0.1, range = 1)
sprbinom(spcov_params_val, data = caribou, xcoord = x, ycoord = y)
sprbinom(spcov_params_val, samples = 5, data = caribou, xcoord = x, ycoord = y)
```

Description

Simulate a spatial gamma random variable with a specific mean and covariance structure.

Usage

```r
sprgamma(
  spcov_params, 
  dispersion = 1, 
  mean = 0, 
  samples = 1, 
  data, 
  randcov_params, 
  partition_factor, 
  ... 
)
```

Arguments

- `spcov_params`: An `spcov_params()` object.
- `dispersion`: The dispersion value.
- `mean`: A numeric vector representing the mean. `mean` must have length 1 (in which case it is recycled) or length equal to the number of rows in `data`. The default is 0.
- `samples`: The number of independent samples to generate. The default is 1.
- `data`: A data frame or `sf` object containing spatial information.
- `randcov_params`: A `randcov_params()` object.
- `partition_factor`: A formula indicating the partition factor.
- `...`: Additional arguments passed to `sprnorm()`.

Details

The values of `spcov_params`, `mean`, and `randcov_params` are assumed to be on the link scale. They are used to simulate a latent normal (Gaussian) response variable using `sprnorm()`. This latent variable is the conditional mean used with dispersion to simulate a gamma random variable.
Value

If samples is 1, a vector of random variables for each row of data is returned. If samples is greater than one, a matrix of random variables is returned, where the rows correspond to each row of data and the columns correspond to independent samples.

Examples

```r
spcov_params_val <- spcov_params("exponential", de = 0.2, ie = 0.1, range = 1)
sprgamma(spcov_params_val, data = caribou, xcoord = x, ycoord = y)
sprgamma(spcov_params_val, samples = 5, data = caribou, xcoord = x, ycoord = y)
```

---

**sprinvgauss**

*Simulate a spatial inverse gaussian random variable*

Description

Simulate a spatial inverse gaussian random variable with a specific mean and covariance structure.

Usage

```r
sprinvgauss(
  spcov_params,
  dispersion = 1,
  mean = 0,
  samples = 1,
  data,
  randcov_params,
  partition_factor,
  ...
)
```

Arguments

- **spcov_params**: An `spcov_params()` object.
- **dispersion**: The dispersion value.
- **mean**: A numeric vector representing the mean. mean must have length 1 (in which case it is recycled) or length equal to the number of rows in data. The default is 0.
- **samples**: The number of independent samples to generate. The default is 1.
- **data**: A data frame or sf object containing spatial information.
- **randcov_params**: A `randcov_params()` object.
- **partition_factor**: A formula indicating the partition factor.
- **...**: Additional arguments passed to `sprnorm()`.
Details
The values of `spcov_params`, `mean`, and `randcov_params` are assumed to be on the link scale. They are used to simulate a latent normal (Gaussian) response variable using `sprnbinom()`. This latent variable is the conditional mean used with dispersion to simulate a inverse gaussian random variable.

Value
If `samples` is 1, a vector of random variables for each row of `data` is returned. If `samples` is greater than one, a matrix of random variables is returned, where the rows correspond to each row of `data` and the columns correspond to independent samples.

Examples
```r
spcov_params_val <- spcov_params("exponential", de = 0.2, ie = 0.1, range = 1)
sprinvgauss(spcov_params_val, data = caribou, xcoord = x, ycoord = y)
sprinvgauss(spcov_params_val, samples = 5, data = caribou, xcoord = x, ycoord = y)
```

**Description**
Simulate a spatial negative binomial random variable with a specific mean and covariance structure.

**Usage**
```r
sprnbinom(
  spcov_params, dispersion = 1, mean = 0, samples = 1, data, randcov_params, partition_factor, ...
)
```

**Arguments**
- `spcov_params`: An `spcov_params()` object.
- `dispersion`: The dispersion value.
- `mean`: A numeric vector representing the mean. `mean` must have length 1 (in which case it is recycled) or length equal to the number of rows in `data`. The default is 0.
- `samples`: The number of independent samples to generate. The default is 1.
data A data frame or `sf` object containing spatial information.
randcov_params A `randcov_params()` object.
partition_factor A formula indicating the partition factor.
... Additional arguments passed to `sprnorm()`.

Details

The values of `spcov_params`, `mean`, and `randcov_params` are assumed to be on the link scale. They are used to simulate a latent normal (Gaussian) response variable using `sprnorm()`. This latent variable is the conditional mean used with dispersion to simulate a negative binomial random variable.

Value

If `samples` is 1, a vector of random variables for each row of `data` is returned. If `samples` is greater than one, a matrix of random variables is returned, where the rows correspond to each row of `data` and the columns correspond to independent samples.

Examples

```r
spcov_params_val <- spcov_params("exponential", de = 0.2, ie = 0.1, range = 1)
sprnbinom(spcov_params_val, data = caribou, xcoord = x, ycoord = y)
sprnbinom(spcov_params_val, samples = 5, data = caribou, xcoord = x, ycoord = y)
```

---

**spnorm**

Simulate a spatial normal (Gaussian) random variable

Description

Simulate a spatial normal (Gaussian) random variable with a specific mean and covariance structure.

Usage

```r
spnorm(  
  spcov_params,  
  mean = 0,  
  samples = 1,  
  data,  
  randcov_params,  
  partition_factor,  
  ...  
)
```

## S3 method for class 'exponential'

```r
spnorm(  
  spcov_params,  
  ...  
)
```
sprnorm

mean = 0,
samples = 1,
data,
randcov_params,
partition_factor,
xcoord,
ycoord,
... )

## S3 method for class 'none'
sprnorm(
  spcov_params,
  mean = 0,
samples = 1,
data,
  randcov_params,
  partition_factor,
  ...
)

## S3 method for class 'car'
sprnorm(
  spcov_params,
  mean = 0,
samples = 1,
data,
  randcov_params,
  partition_factor,
  W,
  row_st = TRUE,
  M,
...
)

Arguments

spcov_params An `spcov_params()` object.
mean A numeric vector representing the mean. mean must have length 1 (in which case it is recycled) or length equal to the number of rows in data. The default is 0.
samples The number of independent samples to generate. The default is 1.
data A data frame or `sf` object containing spatial information.
randcov_params A `randcov_params()` object.
partition_factor A formula indicating the partition factor.
... Other arguments. Not used (needed for generic consistency).
xcoord
Name of the column in data representing the x-coordinate. Can be quoted or unquoted. Not required if data are an sf object.

ycoord
Name of the column in data representing the y-coordinate. Can be quoted or unquoted. Not required if data are an sf object.

W
Weight matrix specifying the neighboring structure used for car and sar models. Not required if data are an sf polygon object and W should be calculated internally (using queen contiguity).

row_st
A logical indicating whether row standardization be performed on W. The default is TRUE.

M
M matrix satisfying the car symmetry condition. The car symmetry condition states that $(I - range * W)^{-1}M$ is symmetric, where $I$ is an identity matrix, range is a constant that controls the spatial dependence, W is the weights matrix, and $^{-1}$ represents the inverse operator. M is required for car models when W is provided and row_st is FALSE. When M, is required, the default is the identity matrix.

Details

Random variables are simulated via the product of the covariance matrix’s square (Cholesky) root and independent standard normal random variables with mean 0 and variance 1. Computing the square root is a significant computational burden and likely unfeasible for sample sizes much past 10,000. Because this square root only needs to be computed once, however, it is nearly the sample computational cost to call sprnorm() for any value of samples.

Only methods for the exponential, none, and car covariance functions are documented here, but methods exist for all other spatial covariance functions defined in spcov_initial(). Syntax for the exponential method is the same as syntax for spherical, gaussian, triangular, circular, cubic, pentaspherical, cosine, wave, jbessel, gravity, rquad, magnetic, matern, cauchy, and pexponential methods. Syntax for the car method is the same as syntax for the sar method. The extra parameter for car and sar models is ignored when all observations have neighbors.

Value

If samples is 1, a vector of random variables for each row of data is returned. If samples is greater than one, a matrix of random variables is returned, where the rows correspond to each row of data and the columns correspond to independent samples.

Examples

```
spcov_params_val <- spcov_params("exponential", de = 1, ie = 1, range = 1)
sprnorm(spcov_params_val, data = caribou, xcoord = x, ycoord = y)
sprnorm(spcov_params_val, mean = 1:30, samples = 5, data = caribou, xcoord = x, ycoord = y)
```
Simulate a spatial Poisson random variable

Description
Simulate a spatial Poisson random variable with a specific mean and covariance structure.

Usage
sprpois(
  spcov_params,
  mean = 0,
  samples = 1,
  data,
  randcov_params,
  partition_factor,
  ...
)

Arguments
spcov_params An `spcov_params()` object.
mean A numeric vector representing the mean. mean must have length 1 (in which case it is recycled) or length equal to the number of rows in data. The default is 0.
samples The number of independent samples to generate. The default is 1.
data A data frame or sf object containing spatial information.
randcov_params A `randcov_params()` object.
partition_factor A formula indicating the partition factor.
... Additional arguments passed to `spnorm()`.

Details
The values of spcov_params, mean, and randcov_params are assumed to be on the link scale. They are used to simulate a latent normal (Gaussian) response variable using `spnorm()`. This latent variable is the conditional mean used with dispersion to simulate a Poisson random variable.

Value
If samples is 1, a vector of random variables for each row of data is returned. If samples is greater than one, a matrix of random variables is returned, where the rows correspond to each row of data and the columns correspond to independent samples.
Examples

```r
spcov_params_val <- spcov_params("exponential", de = 0.2, ie = 0.1, range = 1)
srrpois(spcov_params_val, data = caribou, xcoord = x, ycoord = y)
srrpois(spcov_params_val, samples = 5, data = caribou, xcoord = x, ycoord = y)
```

---

**sulfate**  
*Sulfate atmospheric deposition in the conterminous USA*

---

**Description**

Sulfate atmospheric deposition in the conterminous USA.

**Usage**

```r
sulfate
```

**Format**

An `sf` object with 197 rows and 2 columns.

- **sulfate**: Total wet deposition sulfate in kilograms per hectare.
- **geometry**: `POINT` geometry representing coordinates in a Conus Albers projection (EPSG: 5070).

**Source**

These data were used in the publication listed in References. Data were downloaded from the National Atmospheric Deposition Program National Trends Network.

**References**


---

**sulfate_preds**  
*Locations at which to predict sulfate atmospheric deposition in the conterminous USA*

---

**Description**

Locations at which to predict sulfate atmospheric deposition in the conterminous USA.

**Usage**

```r
sulfate_preds
```
Format

An sf object with 197 rows and 1 column.

- geometry: POINT geometry representing coordinates in a Conus Albers projection (EPSG: 5070).

Source

These data were used in the publication listed in References. Data were downloaded from the National Atmospheric Deposition Program National Trends Network.

References


Description

Summarize a fitted model object.

Usage

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

## S3 method for class 'spautor'
summary(object, ...)

## S3 method for class 'spglm'
summary(object, ...)

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

Arguments

- `object`: A fitted model object from `splm()`, `spautor()`, `spglm()`, or `spgautor()`.
- `...`: Other arguments. Not used (needed for generic consistency).

Details

`summary()` creates a summary of a fitted model object intended to be printed using `print()`. This summary contains useful information like the original function call, residuals, a coefficients table, a pseudo r-squared, and estimated covariance parameters.
Value
A list with several fitted model quantities used to create informative summaries when printing.

See Also
print.spmodel()

Examples
spmod <- splm(z ~ water + tarp,
  data = caribou,
  spcov_type = "exponential", xcoord = x, ycoord = y
)
summary(spmod)

Description
Tidy a fitted model object into a summarized tibble.

Usage
## S3 method for class 'splm'
tidy(x, conf.int = FALSE, conf.level = 0.95, effects = "fixed", ...)

## S3 method for class 'spautor'
tidy(x, conf.int = FALSE, conf.level = 0.95, effects = "fixed", ...)

## S3 method for class 'spglm'
tidy(x, conf.int = FALSE, conf.level = 0.95, effects = "fixed", ...)

## S3 method for class 'spgautor'
tidy(x, conf.int = FALSE, conf.level = 0.95, effects = "fixed", ...)

Arguments
x A fitted model object from splm(), spautor(), spglm(), or spgautor().
conf.int Logical indicating whether or not to include a confidence interval in the tidied output. The default is FALSE.
conf.level The confidence level to use for the confidence interval if conf.int is TRUE. Must be strictly greater than 0 and less than 1. The default is 0.95, which corresponds to a 95 percent confidence interval.
effects The type of effects to tidy. Available options are "fixed" (fixed effects), "spcov" (spatial covariance parameters), and "randcov" (random effect variances). The default is "fixed".
... Other arguments. Not used (needed for generic consistency).
Value

A tidy tibble of summary information effects.

See Also

glance.spmodel() augment.spmodel()

Examples

spmod <- splm(z ~ water + tarp,
  data = caribou,
  spcov_type = "exponential", xcoord = x, ycoord = y
)
tidy(spmod)
tidy(spmod, effects = "spcov")

---

varcomp

Variability component comparison

Description

Compare the proportion of total variability explained by the fixed effects and each variance parameter.

Usage

varcomp(object, ...)

## S3 method for class 'splm'
varcomp(object, ...)

## S3 method for class 'spautor'
varcomp(object, ...)

## S3 method for class 'spglm'
varcomp(object, ...)

## S3 method for class 'spgautor'
varcomp(object, ...)

Arguments

object A fitted model object (e.g., from splm(), spautor(), spglm(), or spgautor()).
...
Other arguments. Not used (needed for generic consistency).
Value

A tibble that partitions the total variability by the fixed effects and each variance parameter. The proportion of variability explained by the fixed effects is the pseudo R-squared obtained by \texttt{psuedoR2()}. The remaining proportion is spread accordingly among each variance parameter: "de", "ie", and if random effects are used, each named random effect. If \texttt{spautor()} objects have unconnected sites, a list is returned with three elements: "connected" for a variability comparison among the connected sites; "unconnected" for a variability comparison among the unconnected sites; and "ratio" for the ratio of the variance of the connected sites relative to the variance of the unconnected sites.

Examples

```r
spmod <- splm(z ~ water + tarp, 
data = caribou, 
  spcov_type = "exponential", xcoord = x, ycoord = y )
varcomp(spmod)
```

---

**vcov.spmodel**

Calculate variance-covariance matrix for a fitted model object

Description

Calculate variance-covariance matrix for a fitted model object.

Usage

```r
## S3 method for class 'splm'
vcov(object, ...)

## S3 method for class 'spautor'
vcov(object, ...)

## S3 method for class 'spglm'
vcov(object, var_correct = TRUE, ...)

## S3 method for class 'spgautor'
vcov(object, var_correct = TRUE, ...)
```

Arguments

- `object`: A fitted model object from \texttt{splm()}, \texttt{spautor()}, \texttt{spglm()}, or \texttt{spgautor()}.
- `...`: Other arguments. Not used (needed for generic consistency).
- `var_correct`: A logical indicating whether to return the corrected variance-covariance matrix for models fit using \texttt{spglm()} or \texttt{spgautor()}. The default is TRUE.
Value

The variance-covariance matrix of coefficients obtained via `coef()`. Currently, only the variance-covariance matrix of the fixed effects is supported.

Examples

```r
spmod <- splm(z ~ water + tarp,
              data = caribou,
              spcov_type = "exponential", xcoord = x, ycoord = y
)
vcov(spmod)
```
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