# Package 'lgpr'

November 21, 2019

Title Longitudinal Gaussian Process Regression

**Version** 0.30.2

Description Implements interpretable nonparametric analysis and covariate selection for longitudinal data using additive
Gaussian process regression. Includes specialized non-stationary disease effect modeling features for biomedical studies. Bayesian inference for model parameters is performed using Stan.

```
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Encoding UTF-8
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Biarch true
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```

VignetteBuilder knitr

2 R topics documented:

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lgpr-package

The 'lgpr' package.

#### **Description**

Longitudinal Gaussian Process regression. The package features

- Additive Gaussian process modeling of longitudinal data
- Posterior inference of the model (hyper)parameters using Stan
- Computation of covariate relevances
- Specialized modeling of a non-stationary disease effect
- Functions for visualizing longitudinal data, posterior samples and model predictions
- Gaussian, Poisson, binomial or negative binomial observation models

#### Basic usage

- See the main function 1gp for creating and fitting additive longitudinal GP models.
- See tutorials at https://jtimonen.github.io/lgpr-usage/

#### Author(s)

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### References

- 1. Carpenter, B. et al. (2017). *Stan: A probabilistic programming language*. Journal of Statistical Software 76(1).
- 2. Jonah Gabry, Ben Goodrich and Martin Lysy (2019). *rstantools: Tools for Developing R Packages Interfacing with 'Stan'*. R package version 2.0.0.
- 3. Gabry, J. and Mahr, T. (2019). *bayesplot: Plotting for Bayesian Models*. R package version 1.7.0, http://mc-stan.org/bayesplot.
- 4. Stan Development Team (2019). *RStan: the R interface to Stan.* R package version 2.19.2. http://mc-stan.org/.

add\_test\_caseIDs

Add case IDs to test data frame

#### **Description**

Add case IDs to test data frame

### Usage

```
add_test_caseIDs(X_test, X_data)
```

6 assess\_convergence

## **Arguments**

X\_test test data frame X\_data data frame

#### Value

Updated X\_test data frame.

affected

Select the affected individuals

### **Description**

Select the affected individuals

### Usage

```
affected(object, medians.return = FALSE, threshold = 0.5)
```

# Arguments

object An object of class lgpfit.

medians.return Should the medians of beta parameters also be returned?

threshold A value that the median of beta has to exceed

#### Value

A binary vector indicating the individuals for which the disease effect is inferred to exist.

assess\_convergence

Assess convergence of the chains

# **Description**

Assess convergence of the chains

# Usage

```
assess_convergence(fit, skip_F_gen = TRUE)
```

# **Arguments**

```
fit An (incomplete) object of class lgpfit.
skip_F_gen Should F_mean, F_var etc. be ignored
```

#### Value

A data frame with columns c("Rhat", "Bulk\_ESS", "Tail\_ESS").

average\_predictions 7

average\_predictions

Average predictions over samples

# Description

Average predictions over samples

# Usage

```
average_predictions(LIST)
```

# Arguments

LIST

a list over samples

# Value

a list

check\_data

Validate the 'data' input to 1gp and resolve covariate types

# Description

Validate the 'data' input to 1gp and resolve covariate types

# Usage

```
check_data(data, varInfo, verbose)
```

# Arguments

data the data frame that was passed to lgp

varInfo variable type info

verbose can this print some info?

# Value

a list

 ${\tt check\_formula}$ 

Validate the formula of 1gp

# Description

Checks if the input 'formula' to lgp\_model are valid with the given data

### Usage

```
check_formula(formula, data)
```

# Arguments

formula the formula that was passed to lgp\_model data the data frame that was passed to lgp\_model

#### Value

nothing

check\_hyperparameter\_names

An error message for wrong hyperparameter naming

# Description

An error message for wrong hyperparameter naming

# Usage

```
check_hyperparameter_names(dist, correct)
```

# Arguments

dist the distribution

correct the allowed hyperparameter names

# Value

nothing

# Description

Component index to covariate index

# Usage

```
component_index_to_covariate_index(D, idx)
```

# Arguments

D integer vector of length 6

idx integer

#### Value

an integer

```
{\tt component\_index\_to\_type}
```

Component index to component type

# Description

Component index to component type

# Usage

```
component_index_to_type(D, idx)
```

# Arguments

D integer vector of length 6

idx integer

# Value

an integer

10 compute\_K\_beta

```
compute_kernel_matrices
```

Evaluate kernel matrices for each component

# Description

Used by compute\_predictions.

# Usage

```
compute_kernel_matrices(X1, X2, kernel_info)
```

### **Arguments**

X1 Covariate matrix of size n1 x sum(D).

X2 Covariate matrix of size n2 x sum(D).

kernel\_info A list of parameters and other kernel info.

#### Value

An array of size  $n1 \times n2 \times sum(D)$ .

 $compute\_K\_beta \qquad \qquad \textit{Compute the multiplier matrix $K\_beta$ (to eneable heterogeneous dis-$ 

ease effect)

# Description

Compute the multiplier matrix K\_beta (to eneable heterogeneous disease effect)

# Usage

```
compute_K_beta(beta, row_to_caseID_1, row_to_caseID_2)
```

# Arguments

```
beta a row vector of length N_cases

row_to_caseID_1

mapping from row index to case ID

row_to_caseID_2

mapping from row index to case ID
```

#### Value

a matrix

compute\_K\_var\_mask 11

compute\_K\_var\_mask

Compute the variance mask kernel matrix

### **Description**

Compute the variance mask kernel matrix

# Usage

```
compute_K_var_mask(disAge1, disAge2, vm_params, stp, nan_replace = 0)
```

# Arguments

disAge1 disease-related age covariate vector of length n1 disAge2 disease-related age covariate vector of length n2

vm\_params vector of two mask function parameters

stp input warping steepness

nan\_replace value to replace nans in disAge vectors

#### Value

a matrix of size n1 x n2

compute\_lppd

Compute log-posterior predictive density at test points

# Description

Compute log-posterior predictive density at test points

# Usage

```
compute_lppd(PRED, y_test)
```

# Arguments

PRED predictions

y\_test values of the response variable at the test points

### Value

```
a matrix with size n_samples x n_data
```

### **Description**

Determine noise level

#### Usage

```
compute_noise_level(pars, model, noise_method)
```

#### **Arguments**

pars A data frame representing one parameter sample, i.e one row of as.data.frame(stanfit),

where stanfit is an object of class stanfit

model An object of class lgpmodel noise\_method Noise level estimation method.

#### Value

a value between 0 and 1

compute\_predicted\_components

Compute component-wise predictions at test points

### **Description**

Used by compute\_predictions.

# Usage

```
compute_predicted_components(KK, KKs, KKss, y_data, sigma_n, DELTA)
```

### **Arguments**

KK Kernel matrices data vs. data.KKs Kernel matrices test vs. data.KKss Kernel matrices test vs. test.

y\_data Response variable.

sigma\_n Noise standard deviation parameter.

DELTA Diagonal jitter that ensures pos. def. kernel.

#### Value

A list containing predicted means and variances.

compute\_predictions 13

compute\_predictions Compute component-wise predictions at test points

### **Description**

Used by lgp\_predict.

#### Usage

```
compute_predictions(X_data, y_data, X_test, params, D, info, cnames, TSCL,
   handle_extra = "warning")
```

### **Arguments**

X\_data Covariate matrix (data points).y\_data Response variable (data points).X\_test Covariate matrix (test points).

params Kernel function and other hyperparameters

D a vector of length 6 info other model info

cnames Names of the model components.

TSCL time scaling function and its inverse

handle\_extra What to do if test data contains individuals that are not in the training data? Must

be 'silent', 'warning' or 'error'.

#### Value

A list.

MCMC sample)

# Description

Compute component relevances and estimate amount of noise (one MCMC sample)

#### Usage

```
compute_relevances(pars, model, method, noise_method)
```

## **Arguments**

pars A data frame representing one parameter sample, i.e one row of as.data.frame(stanfit),

where stanfit is an object of class stanfit

model An object of class lgpmodel

method Relevance determination method. Must be either "f\_mean" or "alpha".

noise\_method Noise level estimation method.

#### Value

```
a matrix of size 1 x n_components + 1
```

compute\_relevances\_alpha

The alpha relevance determination method

### **Description**

The alpha relevance determination method

# Usage

```
compute_relevances_alpha(pars, model)
```

# Arguments

pars A data frame representing one parameter sample, i.e one row of as.data.frame(stanfit),

where stanfit is an object of class stanfit

model An object of class 1gpmodel

#### Value

```
a vector of length n_components
```

compute\_relevances\_fmean

The f\_mean relevance determination method

# Description

The f\_mean relevance determination method

# Usage

```
compute_relevances_fmean(pars, model)
```

### **Arguments**

pars A data frame representing one parameter sample, i.e one row of as.data.frame(stanfit),

where stanfit is an object of class stanfit

model An object of class lgpmodel

#### Value

a vector of length n\_components

create\_covariates\_stan 15

```
create_covariates_stan
```

Create the covariate matrix that is given to stan

# Description

Create the covariate matrix that is given to stan

# Usage

```
create_covariates_stan(data, varInfo, types, formula, verbose)
```

### **Arguments**

data the data frame that was passed to lgp

varInfo original variable type info

types the types returned by check\_data

formula the model formula

verbose can this print some info?

#### Value

a list

### **Description**

A helper function for plot\_data.

### Usage

```
create_data_plot_df(data, hl_1, hl_2, hl_cont)
```

# **Arguments**

data a data frame

hl\_1 highlighting by color
hl\_2 highlighting by linestyle
hl\_cont highlighting continuous

#### Value

an extended data frame

16 create\_F

create\_example\_fit

Create an example fit object

### **Description**

Create an example fit object

#### Usage

```
create_example_fit(N = 4, t = 10 * c(1, 2, 3, 4, 5), iter = 100, chains = 1)
```

#### **Arguments**

N number of individuals

t time points

iter number of iterations chains number of chains

#### Value

an object of class lgpfit

create\_F

Simulate latent function components for longitudinal data analysis

# Description

Simulate latent function components for longitudinal data analysis

### Usage

```
create_F(X, covariates, relevances, lengthscales, X_affected, dis_fun,
  bin_kernel, steepness, vm_params, force_zeromean)
```

## Arguments

Χ

input data matrix (generated by create\_X)

covariates

Integer vector that defines the types of covariates (other than id and age). Different integers correspond to the following covariate types:

- 0 = disease-related age
- 1 = other continuous covariate
- 2 = a categorical covariate that interacts with age
- 3 = a categorical covariate that acts as a group offset
- 4 = a categorical covariate that that acts as a group offset AND is restricted to have value 0 for controls and 1 for cases

relevances Relative relevance of each component. Must have be a vector so that

length(relevances) = 2 + length(covariates).

First two values define the relevance of the infividual-specific age and shared

age component, respectively.

lengthscales A vector so that

length(lengthscales) = 2 + sum(covariates %in% c(0,1,2)).

X\_affected which individuals are affected by the disease

dis\_fun A function or a string that defines the disease effect. If this is a function, that

function is used to generate the effect. If dis\_fun is "gp\_vm" or "gp\_ns", the disease component is drawn from a nonstationary GP prior (vm is the variance

masked version of it).

bin\_kernel Should the binary kernel be used for categorical covariates? If this is TRUE, the

effect will exist only for group 1.

steepness Steepness of the input warping function. This is only used if the disease compo-

nent is in the model.

vm\_params Parameters of the variance mask function. This is only needed if useMaskedVarianceKernel

= TRUE.

force\_zeromean Should each component (excluding the disease age component) be forced to

have a zero mean?

#### Value

a data frame FFF where one column corresponds to one additive data component

create\_predictions\_plot\_df1

Create a plotting data frame for ggplot

# Description

A helper function for plot\_predictions.

#### Usage

```
create_predictions_plot_df1(fit, scale_f = TRUE, n_sds)
```

# Arguments

fit An object of class lgpfit.

scale\_f Should the predictions be scaled back to the original data scale?

n\_sds number of standard deviations for the uncertainty band width

#### Value

a data frame

create\_predictions\_plot\_df2

Create a plotting data frame for ggplot

# Description

A helper function for plot\_predictions.

### Usage

```
create_predictions_plot_df2(model, PRED, scale_f = TRUE, mode, n_sds)
```

# Arguments

model An object of class 1gpmodel.

PRED Predictions computed using lgp\_predict.

scale\_f Should the predictions be scaled back to the original data scale?

mode mode

n\_sds number of standard deviations for the uncertainty band width

### Value

a data frame

create\_simdata\_plot\_df

Create a plotting data frame for ggplot

# Description

A helper function for  $plot\_simdata\_by\_component$ .

# Usage

```
create_simdata_plot_df(simData)
```

# Arguments

simData An object created using simulate\_data.

#### Value

a data frame

create\_stan\_input 19

create_stan_input	Create input for Stan	

### **Description**

Parses the formula and data input to lgp\_model. Also performs many input checks.

# Usage

```
create_stan_input(formula, data, prior, likelihood, varInfo, standardize,
  uncertain_effect_time, equal_effect, C_hat, DELTA, sample_F, verbose,
  variance_mask, N_trials, skip_gen_quant)
```

### **Arguments**

formula	A formula of the form $y \sim x1 + x2 + x3$ defining the response variable y and covariates xi. All variables that appear in the formula must exist as columns of data.
data	A data frame containing (at least) the variables given in formula.
prior	Prior distribution. Can be created for example using the function prior_default.
likelihood	Determines the observation model. Must be either "Gaussian" (default), "Poisson", "NB" (negative binomial) or "binomial".
varInfo	Variable type info.
standardize	Should the response variable be standardized?
uncertain_effe	ct_time
	Do we wish to model uncertainty in the disease effect time?
equal_effect	Is the disease effect assumed to be equally strong for all diseased individuals?
C_hat	This can only be given if likelihood is Poisson or NB. The signal $f$ will the be transformed so that $g = \exp(C_hat + f)$ . If NULL, it will be set to $C_hat = \log(mean(y))$ , where $y$ is the response variable.
DELTA	the amount of added jitter to ensure positive definiteness of the kernel
sample_F	Determines if the function values are be sampled (must be TRUE if likelihood is not Gaussian).
verbose	Should more verbose output be printed?
variance_mask	Should a variance mask be used to force disease component variance to zero before disease onset?
N_trials	This argument (number of trials) is only needed when likelihood is binomial. Must have length one or equal to number of data points. Setting N_trials=1 corresponds to Bernoulli observation model.
skip_gen_quant	If this is true, the generated quantities block of Stan is skipped.

# Value

A list containing the data to be given to rstan::sampling, some info about preprocessing and all the information about scaling the inputs and response, and updated variable type info.

20 create\_X

create\_test\_points

Create a matrix of test points

#### **Description**

Create a matrix of test points

#### Usage

```
create_test_points(object, t_test)
```

### **Arguments**

object An object of class lgpmodel or lgpfit

t\_test Test time points (will be same for each individual).

### Value

A data frame.

create\_X

Simulate an input data frame X

### **Description**

Simulate an input data frame X

# Usage

```
create_X(N, covariates, names, n_categs, t_data, t_jitter, t_effect_range,
    continuous_info, verbose)
```

### **Arguments**

N

Number of individuals.

covariates

Integer vector that defines the types of covariates (other than id and age). If not given, only the id and age covariates are created. Different integers correspond to the following covariate types:

- 0 = disease-related age
- 1 = other continuous covariate
- 2 = a categorical covariate that interacts with age
- 3 = a categorical covariate that acts as a group offset
- 4 = a categorical covariate that that acts as a group offset AND is restricted to have value 0 for controls and 1 for cases

names

Covariate names.

n\_categs

An integer vector defining the number of categories for each categorical covariate, so that length(n\_categs) equals to the number of 2's and 3's in the covariates vector.

create\_X\_star 21

t\_data Measurement times.

t\_jitter Standard deviation of the jitter added to the given measurement times.

t\_effect\_range Time interval from which the disease effect times are sampled uniformly. Alter-

natively, This can any function that returns the (possibly randomly generated)

real disease effect time for one individual.

continuous\_info

Info for generating continuous covariates. Must be a list containing fields lambda and mu, which have length 3. The continuous covariates are generated so that  $x < -\sin(a \times t + b) + c$ , where

• t < -seq(0, 2\*pi, length.out = k)

• a <-mu[1] + lambda[1]\*stats::runif(1)

• b <-mu[2] + lambda[2]\*stats::runif(1)

• c <-mu[3] + lambda[3]\*stats::runif(1)

verbose verbosity mode

#### Value

list(X,onsets,par\_cont)

create\_X\_star

 $Create\ X\_star$ 

# Description

Create X\_star

### Usage

```
create_X_star(X, D, t_test, SCL, X_notnan)
```

#### **Arguments**

X covariate matrix

D covariate type information

t\_test Test time points (will be same for each individual).

SCL time scaling function and its inverse

X\_notnan indicates where X\_diseaseAge is not NaN

### Value

A data frame.

22 disease\_effect

create v	
create v	

Generate noisy observations

# Description

Generate noisy observations

#### Usage

```
create_y(noise_type, f, snr, phi, N_trials)
```

#### **Arguments**

noise\_type Either "Gaussian", "Poisson", NB" (negative binomial) or "binomial".

f The underlying signal.

snr The desired signal-to-noise ratio. This argument is valid only with

noise\_type = "Gaussian".

phi The dispersion parameter for negative binomial data. The variance is  $g + g^2/phi$ .

N\_trials The number of trials parameter for binomial data.

#### Value

A list out, where

- out\$g is f mapped through an inverse link function and
- out\$y is the noisy response variable.

disease\_effect

Draw disease component from a parameteric form

### **Description**

Draw disease component from a parameteric form

# Usage

```
disease_effect(X_id, X_disAge, dis_fun)
```

# Arguments

X\_id the id covariate

X\_disAge the diseaseAge covariate dis\_fun the disease age effect function

# Value

a vector

drawCategorical 23

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drawCatego	ri	cal
uiawcatego	1 1	Сат

Indepedently draw categorical variables for each individual

# Description

Indepedently draw categorical variables for each individual

# Usage

```
drawCategorical(N, k, v)
```

# Arguments

N number of individuals k number of timepoints

v vector of numbers of different categories

# Value

```
a matrix of size N x D, where D <-length(v)
```

 ${\tt drawContinuous}$ 

Indepedently draw continuous variables for each individual

# Description

Indepedently draw continuous variables for each individual

# Usage

```
drawContinuous(N, k, D, mu, lambda)
```

# Arguments

N	number of individuals
k	number of timepoints
D	number of variables
mu	a vector of length 3
lambda	a vector of length 3

## Value

```
a matrix of size N x D
```

24 drawMeasurementTimes

 ${\tt drawLatentComponents} \quad \textit{Draw realizations of multivariate normals}$ 

# Description

Draw realizations of multivariate normals

### Usage

```
drawLatentComponents(KK)
```

# **Arguments**

KK

3D matrix where KK[,,j] is the jth kernel matrix

# Value

a matrix FFF

# Description

Draw the age covariate

### Usage

```
{\tt drawMeasurementTimes}({\tt N}, \ {\tt t\_data}, \ {\tt t\_jitter})
```

# **Arguments**

N number of individuals  $t_{data}$  a vector of length k

t\_jitter Standard deviation of the jitter added to the given measurement times.

# Value

a vector of length N\*k

```
extract_t_effect_samples
```

Extract samples of T\_effect

# Description

Extract samples of T\_effect

# Usage

```
extract_t_effect_samples(fit)
```

# **Arguments**

fit

an object of class lgpfit

# Value

a matrix

full\_model

Create a full model with all covariates included

# Description

Create a full model with all covariates included

# Usage

```
full_model(data, ...)
```

# Arguments

data a data frame

... additional parameters to lgp\_model

# Value

a ggplot object

26 get\_case\_ids

full\_model\_formula

Get formula of a full model with all covariates included

# Description

Get formula of a full model with all covariates included

# Usage

```
full_model_formula(data)
```

# Arguments

data

a data frame, where the response variable is the last column

### Value

a formula

get\_case\_ids

Get case ids in original data

# Description

Get case ids in original data

# Usage

```
get_case_ids(fit)
```

# Arguments

fit

an object of class lgpfit

#### Value

a character vector

get\_case\_row\_mappings Create case ID to rows and back mappings

# Description

Create mappings

- from case ID to data rows (caseID\_to\_rows, caseID\_nrows)
- from row number to case ID (row\_to\_caseID)

### Usage

```
get_case_row_mappings(X_notnan, X_id, only_R2C = FALSE)
```

### **Arguments**

X\_notnan binary vector indicating if diseaseAge is available for that measurement

X\_id the id covariate in X

only\_R2C should this return only the rows-to-caseID mapping

#### Value

a list

get\_diseased\_info

Get some variables related to diseased individuals

### **Description**

Get some variables related to diseased individuals

### Usage

```
get_diseased_info(D, X, X_notnan, uncertain_effect_time, equal_effect,
    TSCL)
```

### **Arguments**

D an integer vector of length 6

X the design matrix

X\_notnan a binary vector of length n

uncertain\_effect\_time

Boolean value

equal\_effect Boolean value

TSCL time scaling function and its inverse

#### Value

a list

 ${\tt get\_function\_components\_from\_df}$ 

Get values of function components at data points, for one MCMC sam-

ple

### **Description**

Get values of function components at data points, for one MCMC sample

# Usage

```
get_function_components_from_df(pars, model)
```

#### **Arguments**

pars A data frame representing one parameter sample, i.e one row of as.data.frame(stanfit),

where stanfit is an object of class stanfit

model An object of class 1gpmodel

#### Value

A matrix of size n\_data x n\_components+2

```
get_function_components_from_df_all
```

Get values of function components at data points

# Description

Get values of function components at data points

#### Usage

```
get_function_components_from_df_all(df, model)
```

### **Arguments**

df A stanfit object as data frame, obtained as as.data.frame(stanfit)

model An object of class lgpmodel

#### Value

An array of size n\_samples x n\_data x n\_components+2

get\_g\_from\_f

 ${\tt get\_g\_from\_f}$ 

 $Get \ signal \ on \ data \ scale \ from \ process \ f$ 

# Description

Get signal on data scale from process f

# Usage

```
get_g_from_f(f, model)
```

# **Arguments**

f A vector

model an object of class 1gpmodel

### Value

A vector g

get\_model\_dims

Get some dimension variables that the Stan model needs as input

# Description

Get some dimension variables that the Stan model needs as input

### Usage

```
get_model_dims(X, D)
```

# Arguments

X the design matrix

D a vector of length 6

# Value

a list

get\_onset\_info

get\_obs\_onset\_times

Extract observed disease onset times from diseaseAge covariate vector

### **Description**

Extract observed disease onset times from diseaseAge covariate vector

### Usage

```
get_obs_onset_times(id, age, disAge)
```

### **Arguments**

id the id covariate, vector of length nage the age covariate, vector of length n

disAge the observed disease-related age covariate, vector of length n

#### Value

vector of observed onset times

get\_onset\_info

Get disease onset info

# Description

This returns

- a vector of observed onsets
- mapping from case ID to average sampling interval before the observed disease onset

# Usage

```
get_onset_info(D, X, MAPS, TSCL)
```

### **Arguments**

D an integer vector of length 6

X the design matrix

MAPS mappings created by get\_case\_row\_mappings

TSCL time scaling function and its inverse

#### Value

two vectors of length N\_cases

get\_pkg\_description 31

get\_pkg\_description

Get lgpr version description

# Description

Get lgpr version description

### Usage

```
get_pkg_description()
```

#### Value

package description

get\_predicted

A helper function

# Description

A helper function

# Usage

```
get_predicted(fit)
```

# **Arguments**

fit

An (incomplete) object of class lgpfit.

# Value

a list

get\_prior\_params

Get prior parameters

# Description

Get prior parameters

# Usage

```
get_prior_params(dist, add_correct)
```

# Arguments

dist the distribution

add\_correct additional correct parameter names

32 get\_response

#### Value

a hyperparameter vector of length 2

get\_prior\_type

A dictionary from distribution names to integer encoding

# Description

A dictionary from distribution names to integer encoding

# Usage

```
get_prior_type(type)
```

# Arguments

type

type of the distribution as a string

#### Value

an integer

get\_response

Get the (scaled) response variable

# Description

Gets and possibly scales the response variable.

# Usage

```
get_response(data, varInfo, standardize, LH)
```

# **Arguments**

data the data frame given as input to 1gp

varInfo variable type info

standardize should the response be standardized to unit variance and zero mean

LH likelihood as integer

# Value

a list with the (scaled) response variable

get\_runtime 33

get\_runtime

Get average runtime of a chain

### **Description**

Get average runtime of a chain

### Usage

```
get_runtime(object)
```

# Arguments

object

An object of class lgpfit.

#### Value

Average runtimes for warmup and sampling

get\_stan\_model

Get main stan model of the package

# Description

Get main stan model of the package

# Usage

```
get_stan_model()
```

# Value

an object of class stanmodel

 ${\tt get\_transform\_type}$ 

A dictionary from transform names to integer encoding

### **Description**

A dictionary from transform names to integer encoding

# Usage

```
get_transform_type(type)
```

### **Arguments**

type

Type of the transform as a string. Allowed arguments are "none" or "square". If NULL, "none" is used.

34 hyperparam\_samples

#### Value

```
an integer (0, 1 or 2)
```

hyperparam\_estimate

Get a posterior estimate of model (hyper)parameters

# Description

Get a posterior estimate of model (hyper)parameters

# Usage

```
hyperparam_estimate(object, type = "mean")
```

### **Arguments**

object An (incomplete) object of class lgpfit.
type Must be "mean", "median", or "map".

### Value

a data frame

hyperparam\_samples

Get a set of model (hyper)parameter samples

# Description

Get a set of model (hyper)parameter samples

# Usage

```
hyperparam_samples(object, samples = NULL)
```

# Arguments

object An (incomplete) object of class lgpfit.

samples Sample indices. If NULL, all samples are taken.

### Value

a data frame

idx\_to\_cont\_index 35

idx\_to\_cont\_index

Component index to how manyth continuous covariate it is

# Description

Component index to how manyth continuous covariate it is

# Usage

```
idx_to_cont_index(D, idx)
```

# **Arguments**

D integer vector of length 6

idx an integer

#### Value

an integer

kernel\_bin

Compute a binary kernel matrix

# Description

Compute a binary kernel matrix

# Usage

```
kernel_bin(x1, x2 = NULL, alpha = 1, pos_class = 1)
```

# Arguments

x1 (integer) vector of length n
 x2 (integer) vector of length m
 alpha marginal std (default = 1)
 pos\_class the positive class label

#### Value

A kernel matrix of size n x m

36 kernel\_se

kernel_ns	Compute a nonstationary kernel matrix using input warping

# Description

Compute a nonstationary kernel matrix using input warping

### Usage

```
kernel_ns(x1, x2 = NULL, alpha = 1, ell, a, b, c, nan_replace = 0)
```

# **Arguments**

x1	vector of length n
x2	vector of length m
alpha	marginal std (default = 1)
ell	lengthscale in the warped space
a	steepness of the warping function rise
b	location of the effective time window
С	maximum range
nan_replace	the value to use for replacing NaN values

# Value

A kernel matrix of size n x m

kernel_se Compute a squared exponential kernel	natrix
--	--------

# Description

Compute a squared exponential kernel matrix

# Usage

```
kernel_se(x1, x2, alpha = 1, ell = 1)
```

# Arguments

x1	vector of length n
x2	vector of length m
alpha	marginal std (default = 1)
ell	lengthscale (default = 1)

### Value

A kernel matrix of size n x m

kernel\_zerosum 37

kerne1_	zerosum

Compute a zeros-sum kernel matrix

## **Description**

Compute a zeros-sum kernel matrix

### Usage

```
kernel\_zerosum(x1, x2, M, alpha = 1)
```

## **Arguments**

x1	(integer) vector of length n
x2	(integer) vector of length m
М	number of categories
alpha	marginal std ( $default = 1$ )

### Value

A (binary) kernel matrix of size n x m

lgp

The main function of the 'lgpr' package

## **Description**

This is a wrapper for both lgp\_model and lgp\_fit. It first creates an lgpmodel object and then fits the model, finally returning an lgpfit object.

# Usage

```
lgp(formula, data, likelihood = "Gaussian", prior = prior_default(),
  uncertain_effect_time = FALSE, equal_effect = TRUE,
  id_variable = "id", time_variable = "age", disAge_variable = NULL,
  continuous_vars = NULL, categorical_vars = NULL,
  offset_vars = NULL, C_hat = NULL, DELTA = 1e-08, sample_F = NULL,
  parallel = FALSE, skip_postproc = FALSE, threshold = 0.95,
  variance_mask = TRUE, N_trials = NULL, relevance_method = "f_mean",
  verbose = FALSE, ...)
```

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## **Arguments**

formula A formula of the form  $y \sim x1 + x2 + x3$  defining the response variable y and

covariates xi. All variables that appear in the formula must exist as columns of

data.

data A data frame containing (at least) the variables given in formula.

likelihood Determines the observation model. Must be either "Gaussian" (default), "Poisson",

"NB" (negative binomial) or "binomial".

prior Prior distribution. Can be created for example using the function prior\_default.

uncertain\_effect\_time

Do we wish to model uncertainty in the disease effect time?

equal\_effect Is the disease effect assumed to be equally strong for all diseased individuals?

id\_variable Name of the unique subject identifier variable.

time\_variable Name of the time variable.

disAge\_variable

Name of the disease-related age variable. If NULL, this will be chosen to be

"diseaseAge", if such covariate is found in the data.

continuous\_vars

Names of other continuous covariates. If NULL, the remaining covariates that

have floating point values are interpreted as continuous.

categorical\_vars

Names of categorical covariates that interact with the time variable. If NULL, the remaining covariates that have integer values are interpreted as categorical.

offset\_vars Names of the categorical covariates that are treated as time-independent group

offsets. If NULL, no variables are interpreted as such covariates.

C\_hat This can only be given if likelihood is Poisson or NB. The signal f will the

be transformed so that  $g = \exp(C_hat + f)$ . If NULL, it will be set to  $C_hat = f$ 

log(mean(y)), where y is the response variable.

DELTA the amount of added jitter to ensure positive definiteness of the kernel

sample\_F Determines if the function values are be sampled (must be TRUE if likelihood is

not Gaussian).

parallel Determines if the chain will be run in parallel (default = FALSE). If TRUE, then

Stan is run by first defining options(mc.cores = parallel::detectCores()).

skip\_postproc In this mode the postprocessing after running Stan is skipped.

threshold Component selection threshold for relevance sum.

variance\_mask Should a variance mask be used to force disease component variance to zero

before disease onset?

N\_trials This argument (number of trials) is only needed when likelihood is binomial.

Must have length one or equal to number of data points. Setting N\_trials=1

corresponds to Bernoulli observation model.

relevance\_method

Component relevance determination method. Must be either "f\_mean" or "alpha".

verbose Should more verbose output be printed?

... Optional arguments passed to rstan::sampling, for example iter, chains or

control. See sampling for the possible arguments.

### Value

An object of class lgpfit.

lgpfit-class 39

lgpfit-class

An S4 class to represent the output of the lgp\_fit function

## **Description**

An S4 class to represent the output of the lgp\_fit function

### **Slots**

```
stan_fit The stanfit object returned by rstan::sampling.
model The lgpmodel object returned by lgp_model.
relevances Inferred component relevances.
selection Component selection info.
pkg_version Package version number.
diagnostics A data frame with columns c("Rhat", "Bulk_ESS", "Tail_ESS").
```

lgpmodel-class

An S4 class to represent an lgp model

### **Description**

An S4 class to represent an lgp model

### **Slots**

```
data The original unmodified data frame.
stan_dat The data to be given as input to rstan::sampling.
scalings Preprocessing scaling functions and their inverse operations.
info Model info.
```

lgp\_component\_names

Get names of model components

# Description

Get names of model components

## Usage

```
lgp_component_names(stan_dat)
```

## **Arguments**

stan\_dat

The data that was passed to rstan::sampling

### Value

names of model components

40 lgp\_fit

lgp\_covariate\_names Get names of model covariates

### **Description**

Get names of model covariates

#### Usage

```
lgp_covariate_names(stan_dat)
```

#### **Arguments**

stan\_dat The data that was passed to rstan::sampling

#### Value

names of model components

lgp\_fit

Fit an lgp model

## **Description**

Samples the posterior of an additive Gaussian process regression model using rstan.

## Usage

```
lgp_fit(model, threshold = 0.95, parallel = FALSE,
    skip_postproc = FALSE, relevance_method = "f_mean",
    verbose = FALSE, ...)
```

# Arguments

model An object of class 1gpmodel.

threshold Component selection threshold for relevance sum.

parallel Determines if the chain will be run in parallel (default = FALSE). If TRUE, then

Stan is run by first defining options (mc.cores = parallel::detectCores()).

skip\_postproc In this mode the postprocessing after running Stan is skipped.

relevance\_method

Component relevance determination method. Must be either "f\_mean" or "alpha".

verbose should some output be printed?

... Optional arguments passed to rstan::sampling, for example iter, chains or

control. See sampling for the possible arguments.

### Value

An object of class lgpfit.

lgp\_model 41

### See Also

For the possible additional arguments, see sampling. For creating the lgpmodel input, see lgp\_model.

lgp_model	Create an lgp model	

## **Description**

Creates an object of class 1gpmode1

### Usage

```
lgp_model(formula, data, likelihood = "Gaussian",
    prior = prior_default(), uncertain_effect_time = FALSE,
    equal_effect = TRUE, C_hat = NULL, DELTA = 1e-08,
    sample_F = NULL, id_variable = "id", time_variable = "age",
    disAge_variable = NULL, continuous_vars = NULL,
    categorical_vars = NULL, offset_vars = NULL, variance_mask = TRUE,
    N_trials = NULL, skip_gen_quant = FALSE, verbose = FALSE)
```

## **Arguments**

formula	A formula of the form $y \sim x1 + x2 + x3$ defining the response variable y and covariates xi. All variables that appear in the formula must exist as columns of data.
data	A data frame containing (at least) the variables given in formula.
likelihood	Determines the observation model. Must be either "Gaussian" (default), "Poisson", "NB" (negative binomial) or "binomial".
prior	Prior distribution. Can be created for example using the function prior_default.
uncertain_effe	ct_time
	Do we wish to model uncertainty in the disease effect time?
equal_effect	Is the disease effect assumed to be equally strong for all diseased individuals?
C_hat	This can only be given if likelihood is Poisson or NB. The signal $f$ will the be transformed so that $g = \exp(C_hat + f)$ . If NULL, it will be set to $C_hat = \log(mean(y))$ , where $y$ is the response variable.
DELTA	the amount of added jitter to ensure positive definiteness of the kernel
sample_F	Determines if the function values are be sampled (must be TRUE if likelihood is not Gaussian).
id_variable	Name of the unique subject identifier variable.
<pre>time_variable disAge_variable</pre>	Name of the time variable.

"diseaseAge", if such covariate is found in the data.

continuous\_vars

Names of other continuous covariates. If NULL, the remaining covariates that have floating point values are interpreted as continuous.

Name of the disease-related age variable. If NULL, this will be chosen to be

42 lgp\_predict

categorical\_vars

Names of categorical covariates that interact with the time variable. If NULL, the

remaining covariates that have integer values are interpreted as categorical.

offset\_vars Names of the categorical covariates that are treated as time-independent group

offsets. If NULL, no variables are interpreted as such covariates.

variance\_mask Should a variance mask be used to force disease component variance to zero

before disease onset?

N\_trials This argument (number of trials) is only needed when likelihood is binomial.

Must have length one or equal to number of data points. Setting N\_trials=1

corresponds to Bernoulli observation model.

skip\_gen\_quant If this is true, the generated quantities block of Stan is skipped.

verbose Should more verbose output be printed?

### Value

An object of class 1gpmodel.

#### See Also

For fitting the model, see lgp\_fit.

lgp\_predict Compute predictions for a fitted model

### **Description**

Compute predictions for a fitted model. Only possible for models with Gaussian likelihood.

### Usage

```
lgp_predict(fit, X_test, samples = "map", print_progress = TRUE,
    print_params = FALSE)
```

# **Arguments**

fit An object of class lgpfit.

X\_test The test points where the predictions should be computed.

samples The predictions can be computed either by using only the posterior mean

(samples="mean"), median (samples="median"), or MAP (samples="map") parameters, or for all parameter samples (samples="all"). This can also be a set of indices, for example samples=c(1:10) gives predictions for the parame-

ter samples 1...10.

print\_progress Should progress be printed (if there is more than one sample)?

print\_params Should the parameter values be printed? (only works if samples is mean or

median.)

## Value

A list.

lgp\_test 43

### See Also

- For creating an lgpfit object, see lgp\_fit.
- For creating an lgpmodel object, see lgp\_model.

lgp\_test

Compute predictions and log-posterior predictive density at test points

# Description

This is a convenience function that wraps lgp\_predict, compute\_lppd and plot\_posterior\_y.

# Usage

```
lgp_test(fit, test_data, plot = FALSE, verbose = TRUE,
  samples = "mean")
```

## **Arguments**

fit an object of class lgpfit

test\_data a test data matrix

should this return also a plot of the data and predictions?

verbose Should this print progress?

samples Sample indices or a keyword "mean", "median", "map", or "all".

## Value

a ggplot object or lppd

likelihood\_as\_int

Convert likelihood string to Stan encoding

## **Description**

Convert likelihood string to Stan encoding

## Usage

```
likelihood_as_int(likelihood)
```

### **Arguments**

```
likelihood a string
```

# Value

an integer

likelihood\_as\_str

Convert the Stan likelihood encoding to a string

# Description

Convert the Stan likelihood encoding to a string

# Usage

```
likelihood_as_str(LH)
```

# Arguments

LH

an integer

# Value

a string

log\_gaussian\_density Compute log-density for gaussian distribution

# Description

Compute log-density for gaussian distribution

# Usage

```
log_gaussian_density(x, mu, s2)
```

# Arguments

x point xmu means2 variance

# Value

a number

matrix\_to\_df 45

matrix\_to\_df

Matrix to data frame without editing column names

# Description

Matrix to data frame without editing column names

# Usage

```
matrix_to_df(M)
```

# Arguments

М

a matrix

# Value

a data frame

model\_info

Get model info

# Description

Get model info

# Usage

```
model_info(object, print = TRUE)
```

# Arguments

object

an object of class lgpmodel or lgpfit

print

should this print the info?

# Value

the info as a string

46 onsetsToDiseaseAge

nameComponents

Create names for all components based on covariate names and types

# Description

Create names for all components based on covariate names and types

# Usage

```
nameComponents(types, names)
```

# Arguments

types vector of covariate types names names of the covariates

# Value

a vector of component names

 $\verb"onsetsToDiseaseAge"$ 

Compute the disease-related ages

# Description

Compute the disease-related ages

# Usage

```
onsetsToDiseaseAge(onsets, age, k)
```

# Arguments

onsets true disease effect times, a vector of length N age the age covariate, a vector of length N\*k k number of measurements per individual

## Value

the diseaseAge covariate, a vector of length N\*k

parse\_prior\_distribution

```
parse_prior_distribution
```

Turn a list describing a prior distribution into vectors to be given to Stan

## **Description**

Turn a list describing a prior distribution into vectors to be given to Stan

## Usage

```
parse_prior_distribution(dist, add_correct = NULL)
```

# **Arguments**

dist a list with field type, and possibly others add\_correct additional correct parameter names

## Value

a list with two vectors to be given to Stan

```
parse_prior_t_effect Turn a list describing an effect time distribution into things to be given to Stan
```

## **Description**

Turn a list describing an effect time distribution into things to be given to Stan

### Usage

```
parse_prior_t_effect(dist, N_cases, T_observed, T_last, UNCRT)
```

## **Arguments**

dist This is prior\$effect, where prior is an argument of lgp\_model

N\_cases number of case individuals

T\_observed observed disease onsets / initiation times
T\_last last time point for each diseased individual

UNCRT 0 or 1

# Value

a list with things to be given to Stan

plot\_beta

```
plot,lgpfit,ANY-method
```

Visualize a fitted 'lgpfit' object

# Description

Visualize a fitted 'lgpfit' object

# Usage

```
## S4 method for signature 'lgpfit,ANY'
plot(fit, x = 1, y = 1, color_scheme = "red")
```

# Arguments

fit an object of class lgpfit

x does nothingy does nothing

color\_scheme bayesplot color scheme

## Value

a ggplot object

plot\_beta

Visualize posterior samples of individual-specific disease effect magnitude parameters

# Description

Can only be used if the disease effect was modeled heterogeneously.

## Usage

```
plot_beta(fit, color_scheme = "red", threshold = 0.5)
```

# **Arguments**

fit An object of class lgpfit.

color\_scheme Name of bayesplot color scheme.

threshold Threshold for median.

#### Value

```
a ggplot object
```

plot\_component 49

plot_component	Helper function for plotting one component	
----------------	--	--

## **Description**

Helper function for plotting one component

## Usage

```
plot_component(MMM, SSS, model, idx, time_is_xvar, linealpha, linetype,
  fill_alpha, X_test, marker, sum_highlight, viridis_option)
```

### **Arguments**

MMM a n array of size n\_samples x n\_data x n\_components
SSS a n array of size n\_samples x n\_data x n\_components

model an object of class 'lgpmodel'
idx Index of component to be plotted.
time\_is\_xvar is the time variable the x-axis variable

linealpha line alpha linetype

fill\_alpha fill alpha for geom\_ribbons
X\_test optional matrix of test points

marker point type

sum\_highlight name of a categorical covariate to be highlighted

viridis\_option the option argument of ggplot2::scale\_colour\_viridis\_c by colour in the

sum plot

### Value

a ggplot object

## **Description**

Helper function for plotting components

## Usage

```
plot_components(MMM, SSS, model, time_is_xvar, X_test = NULL,
    sum_highlight = NULL, linealpha = 1, linetype = 1,
    fill_alpha = 0.3, marker = NULL, ncol = NULL, nrow = NULL,
    legend = NULL, labels = NULL, ylim = NULL, font_size = 9,
    theme = ggplot2::theme_linedraw(), legend_dir = "horizontal",
    xlabel = NULL, ylabel = " ", viridis_option = "viridis",
    return_list = FALSE)
```

#### **Arguments**

MMM a n array of size n\_samples x n\_data x n\_components
SSS a n array of size n\_samples x n\_data x n\_components

model an object of class 'lgpmodel'

time\_is\_xvar is the time variable the x-axis variable

X\_test optional matrix of test points

sum\_highlight name of a categorical covariate to be highlighted

linealpha line alpha
linetype line type

fill\_alpha fill alpha for geom\_ribbons

marker point type

ncol number of plot columns nrow number of plot rows

legend legend argument for ggarrange, use "none" to remove legends

labels labels argument for ggarrange

ylim y axis limits
font\_size font size for plots
theme ggplot theme
legend\_dir direction of legend

xlabel x-axis label ylabel y-axis label

viridis\_option the option argument of ggplot2::scale\_colour\_viridis\_c by colour in the

sum plot

return\_list should this return a list of ggplot objects instead of doing ggarrange

#### Value

an object returned by ggpubr::ggarrange or list

```
plot_components_posterior
```

Visualize inferred components

# Description

Visualize inferred components

## Usage

```
plot_components_posterior(fit, subsamples = NULL, time_is_xvar = TRUE,
    PRED = NULL, marker = NULL, sample_idx = 1, n_sd = 2, ...)
```

#### **Arguments**

fit An object of class lgpfit.

subsamples How many samples to plot. If this is NULL, average over all samples is plotted.

If this is "all", all samples are plotted.

time\_is\_xvar is the time variable the x-axis variable in all subplots?

PRED object returned by lgp\_predict

marker point type

sample\_idx Which sample to plot.

n\_sd number of standard deviations (ribbon width)... additional arguments for plot\_components

### Value

an object returned by ggpubr::ggarrange or a list of ggplot2 objects

plot\_components\_posterior\_sub1

Helperfor plot\_components\_posterior

# Description

Helper for plot\_components\_posterior

#### **Usage**

```
plot_components_posterior_sub1(fit, subsamples, time_is_xvar, marker, ...)
```

## **Arguments**

fit An object of class lgpfit.

subsamples How many samples to plot. If this is NULL, average over all samples is plotted.

If this is "all", all samples are plotted.

time\_is\_xvar is the time variable the x-axis variable in all subplots?

marker point type

... additional arguments for plot\_components

## Value

an object returned by ggpubr::ggarrange or a list

### **Description**

```
Helper for plot_components_posterior
```

#### Usage

## **Arguments**

```
fit An object of class lgpfit.

PRED object returned by lgp_predict
sample_idx Which sample to plot.
```

 $\label{time_is_xvar} \mbox{ is the time variable the $x$-axis variable in all subplots?}$ 

n\_sd number of standard deviations (ribbon width)... additional arguments for plot\_components

#### Value

```
an object returned by ggpubr::ggarrange or a list
```

```
plot_components_simdata
```

Visualize the components of a simulated data set

# Description

Visualize the components of a simulated data set

# Usage

```
plot_components_simdata(simData, time_is_xvar = TRUE, marker = 16, ...)
```

# Arguments

```
simData simulated data object (list)
```

time\_is\_xvar is the time variable the x-axis variable in all subplots?

marker point marker

... additional arguments for plot\_components

## Value

```
an object returned by ggpubr::ggarrange or list
```

plot\_data 53

plot_data	A spaghetti plot of longitudinal data.	

## Description

A spaghetti plot of longitudinal data.

# Usage

```
plot_data(data, highlight = NULL, response = "y", id_variable = "id",
   time_variable = "age", psize = 2, lwd = 0.5, title = NULL)
```

## Arguments

data A data frame.

highlight Name of a covariate to be highlighted with color, or id of a subject to be high-

lighted.

response Name of the response variable.

id\_variable Name of id variable.time\_variable Name of time variable.

psize point size lwd line width

title additional string added to title

### Value

a ggplot object

plot\_data\_hl\_cat A spaghetti plot of longitudinal data, highlighting a categorical covariate.

# Description

A spaghetti plot of longitudinal data, highlighting a categorical covariate.

## Usage

```
plot_data_hl_cat(data, highlight = NULL, response = "y",
   id_variable = "id", time_variable = "age", psize = 2, lwd = 0.5)
```

### **Arguments**

data A data frame.

highlight Name of a categorical covariate to be highlighted with color.

response Name of the response variable.

id\_variable Name of id variable.time\_variable Name of time variable.

psize point size lwd line width

54 plot\_data\_hl\_disease

### Value

a ggplot object

## **Description**

A spaghetti plot of longitudinal data, highlighting a continuous covariate.

## Usage

```
plot_data_hl_cont(data, highlight = NULL, response = "y",
  id_variable = "id", time_variable = "age", psize = 2, lwd = 0.5,
  colgrad = ggplot2::scale_colour_gradient2())
```

### **Arguments**

data A data frame.

highlight Name of a continuous covariate to be highlighted with color.

response Name of the response variable.

id\_variable Name of id variable.time\_variable Name of time variable.

psize point size
lwd line width
colgrad color gradient

### Value

a ggplot object

plot\_data\_hl\_disease A spaghetti plot of longitudinal data, highlighting based on disease group.

# Description

A spaghetti plot of longitudinal data, highlighting based on disease group.

## Usage

```
plot_data_hl_disease(data, highlight = "diseaseAge", response = "y",
   id_variable = "id", time_variable = "age", psize = 2, lwd = 0.5)
```

plot\_data\_hl\_individual

## **Arguments**

data A data frame.

highlight Name of the disease-related age variable.

response Name of the response variable.

id\_variable Name of id variable.time\_variable Name of time variable.

psize point size lwd line width

## Value

a ggplot object

```
plot_data_hl_individual
```

A spaghetti plot of longitudinal data, highlighting one individual.

## **Description**

A spaghetti plot of longitudinal data, highlighting one individual.

## Usage

```
plot_data_hl_individual(data, highlight = 1, response = "y",
   id_variable = "id", time_variable = "age", psize = 2, lwd = 0.5)
```

## **Arguments**

data A data frame.

highlight Number indicating the individual to highlight.

response Name of the response variable.

id\_variable Name of id variable.time\_variable Name of time variable.

psize point size lwd line width

## Value

a ggplot object

plot\_effect\_times

plot\_data\_plain

A spaghetti plot of longitudinal data without highlighting.

# Description

A spaghetti plot of longitudinal data without highlighting.

## Usage

```
plot_data_plain(data, response = "y", id_variable = "id",
    time_variable = "age", psize = 2, lwd = 0.5)
```

## **Arguments**

data A data frame.

response Name of the response variable.

id\_variable Name of id variable.time\_variable Name of time variable.

psize point size lwd line width

### Value

a ggplot object

plot\_effect\_times

Visualize posterior uncertainty in the disease effect times

### **Description**

Can only be used if the uncertainty of effect time was modeled.

## Usage

```
plot_effect_times(fit, color_scheme = "red", prob = 1,
   prob_outer = 1, point_est = "none")
```

## **Arguments**

fit An object of class lgpfit.

color\_scheme Name of bayesplot color scheme.

prob\_outer Outer interval
point\_est Point estimate type

## Value

a ggplot object

plot\_inputwarp 57

plot\_inputwarp

Visualize the input warping function for different parameter samples

# Description

Visualize the input warping function for different parameter samples

### Usage

```
plot_inputwarp(fit, p = 300, color_scheme = "red", b = 0, c = 1)
```

## **Arguments**

fit An object of class 1gpfit.
p number of plot points

color\_scheme Name of bayesplot color scheme.

b location of the effective time window (default = 0)

c maximum range (default = 1)

### Value

a ggplot object

plot\_posterior\_f

*Plot posterior of f* 

# Description

This is a wrapper for plot\_posterior\_predictions.

# Usage

```
plot_posterior_f(fit, PRED = NULL, plot_uncertainty = TRUE,
  data_marker = 16, n_sds = 2, ...)
```

### **Arguments**

fit An object of class lgpfit.

PRED Predictions computed using lgp\_predict.

plot\_uncertainty

Should an uncertainty ribbon be plotted?

data\_marker pch for data points

n\_sdsnumber of standard deviations for the uncertainty band widthadditional arguments to plot\_posterior\_predictions

# Value

```
a ggplot object
```

```
plot_posterior_predictions
```

Plot posterior of f or predictive distribution for y

## **Description**

Plot posterior of f or predictive distribution for y

### Usage

```
plot_posterior_predictions(fit, mode, PRED = NULL,
    color_scheme = "red", color_scheme_t_effect = "gray", alpha = 0.5,
    alpha_line = 1, alpha2 = 0.5, plot_uncertainty = TRUE,
    title = NULL, ylim = NULL, plot_obs_onset = FALSE,
    plot_t_effect_samples = FALSE, ypos_dens = NULL, test_data = NULL,
    color_test = "deepskyblue2", pch_test = 21, size_test = 2,
    error_bar = FALSE, n_sds = 2, reference_times = NULL,
    post_t_effect_stat = "none", original_y_scale = TRUE,
    data_color = "black", data_marker = 21, ons_linetypes = c(1, 2, 3),
    ons_linecolors = c("black", "red", "gray50"))
```

## **Arguments**

```
fit
                  An object of class lgpfit.
                  Must be either "posterior" or "predictive".
mode
PRED
                  Predictions computed using lgp_predict.
color_scheme
                  Name of bayesplot color scheme or a list with fieds 'dark' and 'light'.
color_scheme_t_effect
                  color scheme name for effect time density plotting
                  Ribbon fill opacity.
alpha
alpha_line
                  Line opacity.
alpha2
                  alpha of t_onset density
plot_uncertainty
                  Should an uncertainty ribbon be plotted?
title
                  optional prefix to plot title
                  y axis limits
vlim
plot_obs_onset should the observed disease onset/initiation time be plotted by a vertical line
plot_t_effect_samples
                  should a distribution of sampled effect times be plotted
                  y-position of the density plot
ypos_dens
test_data
                  Test data frame
color_test
                  test point color
pch_test
                  test point marker
size_test
                  test point size
error_bar
                  should uncertainty be plotted using error bars instead of a ribbon
```

plot\_posterior\_y 59

## Value

a ggplot object

plot\_posterior\_y

Plot posterior predictive distribution

## **Description**

This is a wrapper for plot\_posterior\_predictions.

# Usage

```
plot_posterior_y(fit, PRED, uncertainty = "ribbon", test_data = NULL,
  data_marker = 16, n_sds = 2, ...)
```

## Arguments

fit An object of class lgpfit.

PRED Predictions computed using lgp\_predict.

uncertainty Either "none", "ribbon" or "errorbar".

test\_data Test data set.

data\_marker pch for data points

n\_sds number of standard deviations for the uncertainty band width

... additional arguments to plot\_posterior\_predictions

## Value

a ggplot object

```
plot_predictions_add_onsets
```

Add disease onset / effect times to predictions plot

## **Description**

NOTE: currently assumes that diseased individuals come first.

# Usage

```
plot_predictions_add_onsets(fit, h, plot_obs_onset, plot_t_effect_samples,
  idvar, timevar, ypos_dens, color_scheme_t_effect, reference_times,
  post_t_effect_stat, linetypes = c(1, 2, 3), linecolors = c("black",
  "red", "gray50"), alpha2 = 1)
```

# **Arguments**

```
fit
                  An object of class lgpfit.
h
                  a ggplot object
plot_obs_onset a boolean value
plot_t_effect_samples
                  a boolean value
idvar
                  id variable name
timevar
                  time variable name
ypos_dens
                  y position of the estimated onset density
color_scheme_t_effect
                  color scheme
reference_times
                  reference onset times
post_t_effect_stat
                  statistic computed from effect time samples
linetypes
                  onset line types
linecolors
                  onset line colors
alpha2
                  alpha parameter
```

## Value

```
a modified ggplot object
```

```
plot_predictions_options
```

Do input checks and set options for plotting predictions

## **Description**

Do input checks and set options for plotting predictions

## Usage

```
plot_predictions_options(fit, color_scheme, original_y_scale, PRED,
  test_data, color_scheme_t_effect, mode, n_sds)
```

## **Arguments**

fit An object of class lgpfit.
color\_scheme Name of bayesplot color scheme.

original\_y\_scale

Boolean value.

PRED Predictions computed using lgp\_predict.

Another color scheme.

mode mode

n\_sds number of standard deviations for the uncertainty band width

### Value

a list

plot\_relevances

Barplot of covariate relevances

## **Description**

Barplot of covariate relevances

### Usage

```
plot_relevances(object, violin = FALSE, color_scheme = "red", ...)
```

## Arguments

object an object of class lgpfit

violin Should a violin plot be used instead of a boxplot

color\_scheme bayesplot color scheme name

... Additional arguments to ggplot2::geom\_boxplot or ggplot2::geom\_violin.

62 plot\_samples

### Value

a ggplot object

plot\_samples

Visualize the distribution of the model parameter samples

## **Description**

This is a wrapper for functions in the bayesplot package.

## Usage

```
plot_samples(object, pars = character(), regex_pars = character(),
  type = "intervals", prob = 0.5, prob_outer = 0.9,
  color_scheme = "red", point_est = "median", binwidth = NULL,
  transformations = list(), off_diag_args = list(size = 1),
  facet_args = list())
```

## Arguments

object An object of class lgpfit.

pars parameter names

regex\_pars regex for parameter names

type Visualization type. Must be either "dens", "areas", "intervals" (default) or

"hist".

prob\_outer inner interval outer interval

color\_scheme See different color schemes in the bayesplot package.

point\_est the point estimate type

binwidth width of histogram bins if type = "hist"

transformations

the parameter transformations

off\_diag\_args Additional argument list for the pairs plot.

facet\_args additional facetting arguments

## Value

a ggplot object

plot\_simdata 63

plot\_simdata

Plot a simulated longitudinal data set for each individual separately

### **Description**

Plot a simulated longitudinal data set for each individual separately

## Usage

# Arguments

simData a list returned by simulate\_data

linecolor line color

nrow an argument for ggplot2::facet\_wrap
ncol an argument for ggplot2::facet\_wrap

i\_test test point indices
color\_point data point color
color\_test test point color
signal\_name name of signal

y\_transform function to transform the data y

### Value

a ggplot object

## See Also

For plotting each component separately, see  $plot\_components\_simdata$ 

postproc

Finalize the lgpfit object after sampling

# Description

Finalize the lgpfit object after sampling

# Usage

```
postproc(fit, threshold = 0.95, relevance_method = "f_mean",
  verbose = FALSE)
```

64 postproc\_relevances

## **Arguments**

fit An (incomplete) object of class lgpfit.

threshold Threshold for relevance sum. Must be a value between 0 and 1.

relevance\_method

Component relevance determination method. Must be either "f\_mean" or "alpha".

verbose Should some output be printed?

### Value

An updated object of class lgpfit.

postproc\_relevances Compute component relevances and estimate amount of noise (one MCMC sample)

## **Description**

Compute component relevances and estimate amount of noise (one MCMC sample)

## Usage

```
postproc_relevances(fit, relevance_method = "f_mean",
    noise_method = "SSE", verbose = FALSE)
```

# **Arguments**

fit An (incomplete) object of class lgpfit.

 $relevance\_method$ 

Component relevance determination method. Must be either "f\_mean" or "alpha".

noise\_method Noise level determination method. Currently must be "SSE".

verbose Should some output be printed?

### Value

An updated object of class lgpfit.

predict\_preproc 65

predict\_preproc

Preprocess some things before computing predictions

# Description

This is a helper function for lgp\_predict.

# Usage

```
predict_preproc(fit, X_test, samples)
```

# Arguments

fit An object of class lgpfit.

X\_test The test points where the predictions should be computed.

samples The samples argument to lgp\_predict

PRED\_to\_arrays

PRED object to arrays

# Description

PRED object to arrays

# Usage

```
PRED_to_arrays(PRED)
```

# Arguments

PRED

an object returned by lgp\_predict

# Value

a list containing two arrays

66 prior\_LonGP

print\_prior

Human-readable description of a specified prior

## **Description**

Print human-readable info about the prior specification that was used or will be used

## Usage

```
print_prior(object)
```

# Arguments

object

An object of class lgpfit or a valid prior argument for the 'lgp' function.

#### Value

nothing

prior\_default

Create the default prior

# Description

Create the default prior

## Usage

```
prior_default(sigma_alpha = 1)
```

## **Arguments**

sigma\_alpha

Sigma parameter of the student-t distribution for all alpha.

## Value

A list defining a valid prior argument for the 1gp function.

prior\_LonGP

Create a similar default prior as in LonGP (Cheng et. al, 2019)

# Description

Not recommended, because a lengthscale close to 0 is possible.

# Usage

```
prior_LonGP()
```

# Value

A list defining a valid prior argument for the lgp\_model function.

prior\_stan\_to\_readable 67

```
prior_stan_to_readable
```

Human-readable information about the priors in the Stan data object

# **Description**

Human-readable information about the priors in the Stan data object

# Usage

```
prior_stan_to_readable(stan_dat)
```

# **Arguments**

stan\_dat

The list that is passed as data to rstan::sampling.

## Value

Info as a string.

prior\_statement

Human-readable prior statement

# Description

Human-readable prior statement

# Usage

```
prior_statement(parname, TYP, P, dist, row_change = TRUE)
```

# Arguments

parname parameter name
TYP two integers

P three real numbers

dist list of distribution names

row\_change should a newline be last character?

### Value

Sampling statement as a string.

68 repvec

prior\_to\_stan

Get priors as a format that can be input to Stan

## **Description**

Get priors as a format that can be input to Stan

## Usage

```
prior_to_stan(D, prior, HMGNS, UNCRT, N_cases, T_observed, T_last)
```

## **Arguments**

D an integer vector of length 6

prior The prior argument supplied to lgp().

HMGNS Is diseaseAge assumed to have a homogenous effect (1) or not (0)?

UNCRT Boolean value, is uncertainty of disease onset modeled?

N\_cases number of case individuals
T\_observed observed disease onsets

T\_last last time point for each diseased individual

### Value

a list with all things related to priors that Stan needs

repvec

Repeat a vector as a rows of an array

## **Description**

Repeat a vector as a rows of an array

## Usage

```
repvec(v, n)
```

# Arguments

v a vector of length m

n number of times to repeat

# Value

returns an array of size n x m

rtgeom 69

rtgeom	Sample from the 'truncated	l geometric' distribution
--------	----------------------------	---------------------------

# Description

Sample from the 'truncated geometric' distribution

## Usage

```
rtgeom(s, p, n = 1)
```

# Arguments

s an integer

p a number between 0 and 1

n number of samples

## Value

an integer from the interval 1...n

scaleRelevances Scale the effect sizes

# **Description**

Scale the effect sizes

# Usage

```
scaleRelevances(FFF, relevances, force_zeromean, i_skip)
```

## **Arguments**

FFF matrix where one column corresponds to one additive data component

relevances the desired variance of each component (column)

force\_zeromean Should each component (excluding the disease age component) be forced to

have a zero mean.

i\_skip induces of components for which the zero-mean forcing is skipped

### Value

a new matrix FFF

selection

Selection of relevant components

# Description

Selection of relevant components

## Usage

```
selection(object, threshold = 0.95)
```

# Arguments

object An object of class lgpfit.

threshold Threshold for relevance sum. Must be a value between 0 and 1.

### Value

A named list

```
selection_fixed_threshold
```

Select relevant components

# Description

Select relevant components

## Usage

```
selection_fixed_threshold(rel, threshold)
```

# Arguments

rel a named vector of component relevances

threshold value between 0 and 1

### Value

indices of selected components (including "noise" always)

selection\_prob 71

selection_prob	Probabilistic selection of relevant components
----------------	--

## **Description**

Probabilistic selection of relevant components

## Usage

## **Arguments**

object An object of class lgpfit.

p a function defining a density over interval [0,1]

h discretization parameter for computing a quadrature

show\_progbar Should this show a progress bar?

### Value

Selection probabilities for each component

```
selection_prob_fixed_threshold

Selection probabilities using a fixed threshold
```

# Description

Selection probabilities using a fixed threshold

## Usage

```
selection_prob_fixed_threshold(relevances, threshold)
```

# **Arguments**

relevances The relevances\$samples slot of an lgpfit object.

threshold value between 0 and 1

# Value

proportion of times each component was selected

72 set\_C\_hat

 $selection\_prob\_plot$ 

Plot of probabilistic selection of relevant components

# **Description**

Plot of probabilistic selection of relevant components

# Usage

```
selection_prob_plot(PROB, H, P)
```

# **Arguments**

PROB computed probabilities at points H

H a grid on interval [0,1]

P threshold probability distribution evaluated at H

### Value

a ggplot object

set\_C\_hat

Set C\_hat (Poisson and NB observation models)

# Description

Set C\_hat (Poisson and NB observation models)

# Usage

```
set_C_hat(C_hat, response, LH)
```

# Arguments

C\_hat the C\_hat argument given as input to 1gp

response response variable

LH likelihood as int

### Value

a real number

set\_N\_cat 73

set\_N\_cat

 $Count \ numbers \ of \ different \ categories \ for \ each \ categorical \ variable$ 

# Description

Count numbers of different categories for each categorical variable

## Usage

```
set_N_cat(X, D)
```

## Arguments

X the design matrix
D a vector of length 6

#### Value

a numeric vector

set\_N\_trials

*Set N\_trials (binomial and Bernoulli observation models)* 

# Description

Set N\_trials (binomial and Bernoulli observation models)

## Usage

```
set_N_trials(N_trials, response, LH)
```

#### **Arguments**

N\_trials the N\_trials argument given as input to lgp

response response variable

LH likelihood as int

#### Value

a numeric vector

 ${\sf show}, {\sf lgpfit-method}$ 

Show a summary of results of the 1gp function

# Description

Show a summary of results of the 1gp function

## Usage

```
## S4 method for signature 'lgpfit'
show(object)
```

# Arguments

object

an object of class lgpfit

#### Value

nothing

```
show,lgpmodel-method Show a summary of an lgpmodel
```

# Description

Show a summary of an lgpmodel

## Usage

```
## S4 method for signature 'lgpmodel'
show(object)
```

## **Arguments**

object

an object of class lgpmodel

## Value

nothing

```
simdata_colnames_pretty
```

Simulated data column names in a prettier form

#### **Description**

Simulated data column names in a prettier form

#### Usage

```
simdata_colnames_pretty(cn)
```

#### **Arguments**

cn

column names

#### Value

names of model components

simulate\_data

Generate an artificial longitudinal data set

#### **Description**

Generate an artificial longitudinal data set.

#### Usage

```
simulate_data(N, t_data, covariates = c(), names = NULL,
  relevances = c(1, 1, rep(1, length(covariates))), n_categs = rep(2,
  sum(covariates %in% c(2, 3))), t_jitter = 0, lengthscales = rep(12,
  2 + sum(covariates %in% c(0, 1, 2))), f_var = 1,
  noise_type = "Gaussian", snr = 3, phi = 1,
  N_affected = round(N/2), t_effect_range = "auto",
  t_observed = "after_0", C_hat = 0, dis_fun = "gp_vm",
  bin_kernel = FALSE, steepness = 0.5, vm_params = c(0.025, 1),
  continuous_info = list(mu = c(pi/8, pi, -0.5), lambda = c(pi/8, pi,
  1)), N_trials = 1, verbose = FALSE, force_zeromean = TRUE)
```

#### **Arguments**

N Number of individuals.

t\_data Measurement times.

covariates

Integer vector that defines the types of covariates (other than id and age). If not given, only the id and age covariates are created. Different integers correspond to the following covariate types:

- 0 = disease-related age
- 1 = other continuous covariate

76 simulate\_data

• 2 = a categorical covariate that interacts with age

- 3 = a categorical covariate that acts as a group offset
- 4 = a categorical covariate that that acts as a group offset AND is restricted to have value 0 for controls and 1 for cases

names Covariate names.

relevances Relative relevance of each component. Must have be a vector so that

length(relevances) = 2 + length(covariates).

First two values define the relevance of the infividual-specific age and shared

age component, respectively.

n\_categs An integer vector defining the number of categories for each categorical co-

variate, so that length(n\_categs) equals to the number of 2's and 3's in the

covariates vector.

t\_jitter Standard deviation of the jitter added to the given measurement times.

lengthscales A vector so that

length(lengthscales) = 2 + sum(covariates %in% c(0,1,2)).

f\_var variance of f

noise\_type Either "Gaussian", "Poisson", NB" (negative binomial) or "binomial".

snr The desired signal-to-noise ratio. This argument is valid only with

noise\_type = "Gaussian".

phi The dispersion parameter for negative binomial data. The variance is  $g + g^2/phi$ .

N\_affected Number of diseased individuals that are affected by the disease. This defaults

to the number of diseased individuals. This argument can only be given if

covariates contains a zero.

t\_effect\_range Time interval from which the disease effect times are sampled uniformly. Alter-

natively, This can any function that returns the (possibly randomly generated)

real disease effect time for one individual.

t\_observed Determines how the disease effect time is observed. This can be any function

that takes the real disease effect time as an argument and returns the (possibly randomly generated) observed onset/initiation time. Alternatively, this can be a

string of the form "after\_n" or "random\_p" or "exact".

C\_hat A constant added to f

dis\_fun A function or a string that defines the disease effect. If this is a function, that

function is used to generate the effect. If dis\_fun is "gp\_vm" or "gp\_ns", the disease component is drawn from a nonstationary GP prior (vm is the variance

masked version of it).

bin\_kernel Should the binary kernel be used for categorical covariates? If this is TRUE, the

effect will exist only for group 1.

steepness Steepness of the input warping function. This is only used if the disease compo-

nent is in the model.

vm\_params Parameters of the variance mask function. This is only needed if useMaskedVarianceKernel

= TRUE.

continuous\_info

Info for generating continuous covariates. Must be a list containing fields lambda and mu, which have length 3. The continuous covariates are generated so that  $x < -\sin(a \times t + b) + c$ , where

• t <-seq(0,2\*pi,length.out = k)

simulate\_kernels 77

```
a <-mu[1] + lambda[1]*stats::runif(1)</li>
b <-mu[2] + lambda[2]*stats::runif(1)</li>
c <-mu[3] + lambda[3]*stats::runif(1)</li>

The number of trials parameter for binomial data.
verbosity mode
```

verbose verbosity mode

force\_zeromean Should each component (excluding the disease age component) be forced to have a zero mean?

#### Value

A list out, where

N\_trials

- out\$data is a data frame containing the actual data and
- out\$components contains more points for smoother visualizations of the generating process.
- out\$onsets contains the real disease effect times
- out\$p\_signal proportion of variance explained by signal

#### **Examples**

```
# Generate Gaussian data
dat <- simulate_data(N = 4, t_data = c(6,12,24,36,48), snr = 3)
# Generate negative binomially distributed count data
dat <- simulate_data(N = 6, t_data = seq(2, 10, by = 2), noise_type = "NB", phi = 2)</pre>
```

simulate\_kernels

Compute all kernel matrices when simulating data

## Description

Compute all kernel matrices when simulating data

## Usage

```
simulate_kernels(X, types, lengthscales, X_affected, bin_kernel,
  useMaskedVarianceKernel, steepness, vm_params)
```

#### **Arguments**

X covariates types vector of covariate types, so that

• 1 = ID

• 2 = age

• 3 = diseaseAge

• 4 = other continuous covariate

• 5 = a categorical covariate that interacts with age

• 6 = a categorical covariate that acts as an offset

lengthscales vector of lengthscales

78 sim\_data\_to\_observed

X\_affected which individuals are affected by the disease

bin\_kernel whether or not binary (mask) kernel should be used for categorical covariates

useMaskedVarianceKernel

should the masked variance kernel be used for drawing the disease component

steepness steepness of the input warping function
vm\_params parameters of the variance mask function

#### Value

a 3D array

#### **Description**

Input check for the covariates-related arguments of simulate\_data

#### Usage

```
sim_check_covariates(covariates, relevances, names, n_cat)
```

#### **Arguments**

covariates argument to simulate\_data
relevances argument to simulate\_data
names argument to simulate\_data

n\_cat the n\_categs argument to simulate\_data

#### Value

the covariate names

sim\_data\_to\_observed Real generated disease ages to observed ones

#### **Description**

Real generated disease ages to observed ones

#### Usage

```
sim_data_to_observed(dat, t_observed)
```

#### **Arguments**

data frame

t\_observed Determines how the disease onset is observed. See documentation of simulate\_data.

#### Value

a new data frame and observed onsets

sim\_generate\_names 79

sim\_generate\_names

Generate names for covariates

# Description

Generate names for covariates

# Usage

```
sim_generate_names(covariates)
```

# Arguments

covariates

vector of covariate types

#### Value

covariate names

sim\_parse\_t\_obs

Parse the t\_observed argument of simulate\_data

# Description

Parse the t\_observed argument of  $simulate\_data$ 

# Usage

```
\\ sim\_parse\_t\_obs(t\_observed)
```

# Arguments

t\_observed a string

#### Value

a list with a name and number

split\_data\_by\_id

split\_data

Split data into training and test data according to given row indices

## **Description**

Split data into training and test data according to given row indices

## Usage

```
split_data(data, i_test, sort_ids = TRUE)
```

## **Arguments**

data a data frame

i\_test test data row indices

sort\_ids should the test indices be sorted into increasing order

#### Value

```
a list(train, test)
```

split\_data\_by\_id

Split data into training and test data according to given individuals

# Description

Split data into training and test data according to given individuals

## Usage

```
split_data_by_id(data, test_ids, id_variable = "id")
```

# Arguments

data a data frame

test\_ids test data individual identifiers

id\_variable name of id variable

## Value

```
alist(train, test)
```

```
{\tt split\_data\_by\_timepoint}
```

Split data into training and test data according to time point indices

## Description

Split data into training and test data according to time point indices

#### Usage

```
split_data_by_timepoint(data, test_idx, id_variable = "id",
   time_variable = "age")
```

#### **Arguments**

data a data frame

#### Value

```
alist(train,test)
```

split\_data\_random

Split data into training and test data randomly

#### **Description**

Split data into training and test data randomly

## Usage

```
split_data_random(data, p_test = 0.1, n_test = NULL)
```

#### **Arguments**

data a data frame

p\_test desired proportion of test data

n\_test desired number of test data points (if NULL, p\_test is used to compute this)

#### Value

```
a list(train, test)
```

82 standardize\_inputs

```
split_data_random_each
```

Split data into training and test data by selecting randomly k points from each individual

## Description

Split data into training and test data by selecting randomly k points from each individual

#### Usage

```
split_data_random_each(data, n_test = 1, id_variable = "id",
   time_variable = "age")
```

#### Arguments

data a data frame

n\_test desired number of test data points per individual

id\_variable name of id variable
time\_variable name of time variable

#### Value

```
alist(train, test)
```

standardize\_inputs

Standardize continuous input variables in X

# Description

Standardize continuous input variables in X

#### Usage

```
standardize_inputs(X, D)
```

## Arguments

X the design matrix

D the covariate types, a vector of length 6

#### Value

updated X and info about scaling

stan\_input\_X\_and\_D 83

#### **Description**

Reorders covariates and takes only those that are needed

#### Usage

```
stan_input_X_and_D(data, varInfo, types, formula, verbose)
```

#### **Arguments**

data a data frame containing the covariates

varInfo original variable type info types types of the covariates

formula model formula

verbose can this print some info?

#### Value

X and needed types and updated varInfo

validate\_prior Validate prior by sampling the signal and noise from it

## Description

Validate prior by sampling the signal and noise from it

# Usage

```
validate_prior(model, chains = 4, iter = 1000, parallel = FALSE)
```

# Arguments

model An object of class lgpmodel.

chains how many chains are used to sample from the prior

iter for how many iterations are the chains run parallel should the chains be run in parallel?

#### Value

An object of class lgpfit and random samples of both 'f' and 'y'.

84 warp\_input

warp\_input

Warp inputs

# Description

Warp inputs

# Usage

```
warp_input(t, a, b, c)
```

# **Arguments**

t a vector

a steepness of the rise

b location of the effective time window

c maximum range

# Value

```
a vector of warped inputs w(t)
```

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