

Package ‘lgpr’

October 31, 2025

Title Longitudinal Gaussian Process Regression

Version 1.2.5

Description Interpretable nonparametric modeling of longitudinal data using additive Gaussian process regression. Contains functionality for inferring covariate effects and assessing covariate relevances. Models are specified using a convenient formula syntax, and can include shared, group-specific, non-stationary, heterogeneous and temporally uncertain effects. Bayesian inference for model parameters is performed using 'Stan'. The modeling approach and methods are described in detail in Timonen et al. (2021) <[doi:10.1093/bioinformatics/btab021](https://doi.org/10.1093/bioinformatics/btab021)>.

License GPL (>= 3)

Encoding UTF-8

LazyData true

Biarch true

Depends R (>= 3.4.0), methods

Imports Rcpp (>= 0.12.0), RcppParallel (>= 5.0.2), RCurl (>= 1.98), rstan (>= 2.26.0), rstantools (>= 2.3.1), bayesplot (>= 1.7.0), MASS (>= 7.3-50), stats (>= 3.4), ggplot2 (>= 3.1.0), gridExtra (>= 0.3.0)

LinkingTo BH (>= 1.75.0-0), Rcpp (>= 1.0.6), RcppEigen (>= 0.3.3.9.1), RcppParallel (>= 5.0.2), rstan (>= 2.26.0), StanHeaders (>= 2.26.0)

SystemRequirements GNU make

NeedsCompilation yes

RoxygenNote 7.3.2

Suggests knitr, rmarkdown, testthat, covr

URL <https://github.com/jtimonen/lgpr>

BugReports <https://github.com/jtimonen/lgpr/issues>

VignetteBuilder knitr

Author Juho Timonen [aut, cre] (ORCID:
<https://orcid.org/0000-0003-2341-6765>),
 Andrew Johnson [ctb]

Maintainer Juho Timonen <juho.timonen@iki.fi>

Repository CRAN

Date/Publication 2025-10-30 23:50:14 UTC

Contents

lgpr-package	3
add_dis_age	5
add_factor	6
add_factor_crossing	6
adjusted_c_hat	7
apply_scaling	8
as_character	8
create_model	9
create_model.covs_and_comps	10
create_model.formula	11
create_model.likelihood	12
create_model.options	13
create_model.prior	14
create_plot_df	15
create_scaling	15
dinvgamma_stanlike	16
draw_pred	16
example_fit	17
fit_summary	18
GaussianPrediction-class	18
get_draws	19
get_pred	20
kernel	20
KernelComputer-class	22
lgp	23
lgpexpr-class	26
lgpfit-class	27
lgpformula-class	28
lgpmodel-class	29
lgprhs-class	30
lgpscaling-class	30
lgpsim-class	31
lgpterm-class	32
model_summary	32
new_x	33
operations	34
plot_api_c	34
plot_api_g	36

plot_components	37
plot_data	39
plot_draws	40
plot_inputwarp	41
plot_invgamma	42
plot_pred	43
plot_sim	44
ppc	45
pred	46
Prediction-class	47
priors	48
prior_pred	49
prior_to_num	50
read_proteomics_data	51
relevances	52
s4_generics	52
sample_model	54
select	55
show	56
sim.create_f	57
sim.create_x	58
sim.create_y	59
sim.kernels	60
simulate_data	61
split	63
testdata_001	64
testdata_002	65
validate	66
var_mask	66
warp_input	67
Index	68

lgpr-package

The 'lgpr' package.

Description

Interpretable nonparametric modeling of longitudinal data using additive Gaussian process regression. Contains functionality for inferring covariate effects and assessing covariate relevances. Models are specified using a convenient formula syntax, and can include shared, group-specific, non-stationary, heterogeneous and temporally uncertain effects. Bayesian inference for model parameters is performed using 'Stan' ([rstan](#)). The modeling approach and methods are described in detail in [Timonen et al. \(2021\)](#).

Core functions

Main functionality of the package consists of creating and fitting an additive GP model:

- `lgp`: Specify and fit an additive GP model with one command.
- `create_model`: Define an `lgpmodel` object.
- `sample_model`: Fit a model by sampling the posterior distribution of its parameters and create an `lgpfit` object.
- `pred`: Computing model predictions and inferred covariate effects after fitting a model.
- `relevances`: Assessing covariate relevances after fitting a model.
- `prior_pred`: Prior predictive sampling to check if your prior makes sense.

Visualization

- `plot_pred`: Plot model predictions.
- `plot_components`: Visualize inferred model components.
- `plot_draws`: Visualize parameter draws.
- `plot_data`: Visualize longitudinal data.

Data

The data that you wish to analyze with 'lgpr' should be in an R `data.frame` where columns correspond to measured variables and rows correspond to observations. Some functions that can help working with such data frames are:

- `new_x`: Creating new test points where the posterior distribution of any function component or sum of all components, or the posterior predictive distribution can be computed after model fitting.
- Other functions: `add_factor`, `add_factor_crossing`, `add_dis_age`, `adjusted_c_hat`.

Vignettes and tutorials

See <https://jtimonen.github.io/lgpr-usage/index.html>. The tutorials focus on code and use cases, whereas the [Mathematical description of lgpr models](#) vignette describes the statistical models and how they can be customized in 'lgpr'.

Citation

Run `citation("lgpr")` to get citation information.

Feedback

Bug reports, PRs, enhancement ideas or user experiences in general are welcome and appreciated. Create an issue in Github or email the author.

Author(s)

Juho Timonen (first.last at iki.fi)

References

1. Timonen, J. et al. (2021). *lgpr: an interpretable non-parametric method for inferring covariate effects from longitudinal data*. Bioinformatics, [url](#).
2. Carpenter, B. et al. (2017). *Stan: A probabilistic programming language*. Journal of Statistical Software 76(1).

See Also

Useful links:

- <https://github.com/jtimonen/lgpr>
- Report bugs at <https://github.com/jtimonen/lgpr/issues>

add_dis_age

Easily add the disease-related age variable to a data frame

Description

Creates the disease-related age covariate vector based on the disease initiation times and adds it to the data frame

Usage

```
add_dis_age(data, t_init, id_var = "id", time_var = "age")
```

Arguments

<code>data</code>	the original data frame
<code>t_init</code>	A named vector containing the observed initiation or onset time for each individual. The names, i.e. <code>names(t_init)</code> , should specify the individual id.
<code>id_var</code>	name of the id variable in data
<code>time_var</code>	name of the time variable in data

Value

A data frame with one column added. The new column will be called `dis_age`. For controls, its value will be `NaN`.

See Also

Other data frame handling functions: [add_factor\(\)](#), [add_factor_crossing\(\)](#), [adjusted_c_hat\(\)](#), [new_x\(\)](#), [split\(\)](#)

add_factor	<i>Easily add a categorical covariate to a data frame</i>
------------	---

Description

Easily add a categorical covariate to a data frame

Usage

```
add_factor(data, x, id_var = "id")
```

Arguments

data	the original data frame
x	A named vector containing the category for each individual. The names should specify the individual id.
id_var	name of the id variable in data

Value

A data frame with one column added. The new column will have same name as the variable passed as input x.

See Also

Other data frame handling functions: [add_dis_age\(\)](#), [add_factor_crossing\(\)](#), [adjusted_c_hat\(\)](#), [new_x\(\)](#), [split\(\)](#)

add_factor_crossing	<i>Add a crossing of two factors to a data frame</i>
---------------------	--

Description

Add a crossing of two factors to a data frame

Usage

```
add_factor_crossing(data, fac1, fac2, new_name)
```

Arguments

data	a data frame
fac1	name of first factor, must be found in df
fac2	name of second factor, must be found in df
new_name	name of the new factor

Value

a data frame

See Also

Other data frame handling functions: [add_dis_age\(\)](#), [add_factor\(\)](#), [adjusted_c_hat\(\)](#), [new_x\(\)](#), [split\(\)](#)

adjusted_c_hat	<i>Set the GP mean vector, taking TMM or other normalization into account</i>
----------------	---

Description

Creates the `c_hat` input for `lgp`, so that it accounts for normalization between data points in the "poisson" or "nb" observation model

Usage

```
adjusted_c_hat(y, norm_factors)
```

Arguments

<code>y</code>	response variable, vector of length <code>n</code>
<code>norm_factors</code>	normalization factors, vector of length <code>n</code>

Value

a vector of length `n`, which can be used as the `c_hat` input to the `lgp` function

See Also

Other data frame handling functions: [add_dis_age\(\)](#), [add_factor\(\)](#), [add_factor_crossing\(\)](#), [new_x\(\)](#), [split\(\)](#)

apply_scaling	<i>Apply variable scaling</i>
---------------	-------------------------------

Description

Apply variable scaling

Usage

```
apply_scaling(scaling, x, inverse = FALSE)
```

Arguments

scaling	an object of class lgpscaling
x	object to which apply the scaling (numeric)
inverse	whether scaling should be done in inverse direction

Value

a similar object as x

See Also

Other variable scaling functions: [create_scaling\(\)](#)

as_character	<i>Character representations of different formula objects</i>
--------------	---

Description

Character representations of different formula objects

Usage

```
## S4 method for signature 'lgpexpr'
as.character(x)

## S4 method for signature 'lgpterm'
as.character(x)

## S4 method for signature 'lgpformula'
as.character(x)
```

Arguments

x	an object of some S4 class
---	----------------------------

Value

a character representation of the object

create_model	Create a model
--------------	----------------

Description

See the [Mathematical description of lgpr models](#) vignette for more information about the connection between different options and the created statistical model.

Usage

```
create_model(
  formula,
  data,
  likelihood = "gaussian",
  prior = NULL,
  c_hat = NULL,
  num_trials = NULL,
  options = NULL,
  prior_only = FALSE,
  verbose = FALSE,
  sample_f = !(likelihood == "gaussian")
)
```

Arguments

formula	The model formula, where • it must contain exactly one tilde (~), with response variable on the left-hand side and model terms on the right-hand side • terms are separated by a plus (+) sign • all variables appearing in formula must be found in data See the "Model formula syntax" section below (lgp) for instructions on how to specify the model terms.
data	A <code>data.frame</code> where each column corresponds to one variable, and each row is one observation. Continuous covariates and the response variable must have type "numeric" and categorical covariates must have type "factor". Missing values should be indicated with <code>NaN</code> or <code>NA</code>. The response variable cannot contain missing values. Column names should not contain trailing or leading underscores.
likelihood	Determines the observation model. Must be either "gaussian" (default), "poisson", "nb" (negative binomial), "binomial" or "bb" (beta binomial).
prior	A named list, defining the prior distribution of model (hyper)parameters. See the "Defining priors" section below (lgp).

c_hat	The GP mean. This should only be given if <code>sample_f</code> is TRUE, otherwise the GP will always have zero mean. If <code>sample_f</code> is TRUE, the given <code>c_hat</code> can be a vector of length <code>dim(data)[1]</code>, or a real number defining a constant GP mean. If not specified and <code>sample_f</code> is TRUE, <code>c_hat</code> is set to • <code>c_hat = mean(y)</code>, if likelihood is "gaussian", • <code>c_hat = log(mean(y))</code> if likelihood is "poisson" or "nb", • <code>c_hat = log(p/(1-p))</code>, where $p = \text{mean}(y/\text{num_trials})$ if likelihood is "binomial" or "bb", where <code>y</code> denotes the response variable measurements.
num_trials	This argument (number of trials) is only needed when likelihood is "binomial" or "bb". Must have length one or equal to the number of data points. Setting <code>num_trials=1</code> and <code>likelihood="binomial"</code> corresponds to Bernoulli observation model.
options	A named list with the following possible fields: • <code>delta</code> Amount of added jitter to ensure positive definite covariance matrices. • <code>vm_params</code> Variance mask function parameters (numeric vector of length 2). If <code>options</code> is NULL, default options are used. The defaults are equivalent to <code>options = list(delta = 1e-8, vm_params = c(0.025, 1))</code>.
prior_only	Should likelihood be ignored? See also sample_param_prior which can be used for any lgpmodel, and whose runtime is independent of the number of observations.
verbose	Should some informative messages be printed?
sample_f	Determines if the latent function values are sampled (must be TRUE if likelihood is not "gaussian"). If this is TRUE, the response variable will be normalized to have zero mean and unit variance.

Value

An object of class [lgpmodel](#), containing the Stan input created based on parsing the specified formula, prior, and other options.

See Also

Other main functions: [draw_pred\(\)](#), [get_draws\(\)](#), [lgp\(\)](#), [pred\(\)](#), [prior_pred\(\)](#), [sample_model\(\)](#)

`create_model.covs_and_comps`

Parse the covariates and model components from given data and formula

Description

Parse the covariates and model components from given data and formula

Usage

```
create_model.covs_and_comps(data, model_formula, x_cont_scl, verbose)
```

Arguments

data	A data.frame where each column corresponds to one variable, and each row is one observation. Continuous covariates and the response variable must have type "numeric" and categorical covariates must have type "factor". Missing values should be indicated with NaN or NA. The response variable cannot contain missing values. Column names should not contain trailing or leading under-scores.
model_formula	an object of class lgpformula
x_cont_scl	Information on how to scale the continuous covariates. This can either be • an existing list of objects with class lgpscaling, or • NA, in which case such list is created by computing mean and standard deviation from data
verbose	Should some informative messages be printed?

Value

parsed input to Stan and covariate scaling, and other info

See Also

Other internal model creation functions: [create_model.formula\(\)](#), [create_model.likelihood\(\)](#), [create_model.prior\(\)](#)

`create_model.formula` *Create a model formula*

Description

Checks if formula is in advanced format and translates if not.

Usage

```
create_model.formula(formula, data, verbose = FALSE)
```

Arguments

formula	The model formula, where • it must contain exactly one tilde (~), with response variable on the left-hand side and model terms on the right-hand side • terms are separated by a plus (+) sign • all variables appearing in formula must be found in data
---------	---

	See the "Model formula syntax" section below (lgp) for instructions on how to specify the model terms.
data	A data.frame where each column corresponds to one variable, and each row is one observation. Continuous covariates and the response variable must have type "numeric" and categorical covariates must have type "factor". Missing values should be indicated with NaN or NA. The response variable cannot contain missing values. Column names should not contain trailing or leading under-scores.
verbose	Should some informative messages be printed?

Value

an object of class [lgpformula](#)

See Also

Other internal model creation functions: [create_model.covs_and_comps\(\)](#), [create_model.likelihood\(\)](#), [create_model.prior\(\)](#)

`create_model.likelihood`

Parse the response variable and its likelihood model

Description

Parse the response variable and its likelihood model

Usage

```
create_model.likelihood(
  data,
  likelihood,
  c_hat,
  num_trials,
  y_name,
  sample_f,
  verbose
)
```

Arguments

data	A data.frame where each column corresponds to one variable, and each row is one observation. Continuous covariates and the response variable must have type "numeric" and categorical covariates must have type "factor". Missing values should be indicated with NaN or NA. The response variable cannot contain missing values. Column names should not contain trailing or leading under-scores.
------	---

likelihood	Determines the observation model. Must be either "gaussian" (default), "poisson", "nb" (negative binomial), "binomial" or "bb" (beta binomial).
c_hat	The GP mean. This should only be given if sample_f is TRUE, otherwise the GP will always have zero mean. If sample_f is TRUE, the given c_hat can be a vector of length <code>dim(data)[1]</code>, or a real number defining a constant GP mean. If not specified and sample_f is TRUE, c_hat is set to • <code>c_hat = mean(y)</code>, if likelihood is "gaussian", • <code>c_hat = log(mean(y))</code> if likelihood is "poisson" or "nb", • <code>c_hat = log(p/(1-p))</code>, where <code>p = mean(y/num_trials)</code> if likelihood is "binomial" or "bb", where y denotes the response variable measurements.
num_trials	This argument (number of trials) is only needed when likelihood is "binomial" or "bb". Must have length one or equal to the number of data points. Setting <code>num_trials=1</code> and <code>likelihood="binomial"</code> corresponds to Bernoulli observation model.
y_name	Name of response variable
sample_f	Determines if the latent function values are sampled (must be TRUE if likelihood is not "gaussian"). If this is TRUE, the response variable will be normalized to have zero mean and unit variance.
verbose	Should some informative messages be printed?

Value

a list of parsed options

See Also

Other internal model creation functions: `create_model.covs_and_comps()`, `create_model.formula()`, `create_model.prior()`

`create_model.options` *Parse the given modeling options*

Description

Parse the given modeling options

Usage

```
create_model.options(options, verbose)
```

Arguments

- | | |
|---------|--|
| options | <p>A named list with the following possible fields:</p> <ul style="list-style-type: none"> • delta Amount of added jitter to ensure positive definite covariance matrices. • vm_params Variance mask function parameters (numeric vector of length 2). <p>If options is NULL, default options are used. The defaults are equivalent to <code>options = list(delta = 1e-8, vm_params = c(0.025, 1))</code>.</p> |
| verbose | Should some informative messages be printed? |

Value

a named list of parsed options

create_model.prior	<i>Parse given prior</i>
--------------------	--------------------------

Description

Parse given prior

Usage

```
create_model.prior(prior, stan_input, verbose)
```

Arguments

- | | |
|------------|--|
| prior | A named list, defining the prior distribution of model (hyper)parameters. See the "Defining priors" section below (lgp). |
| stan_input | a list of stan input fields |
| verbose | Should some informative messages be printed? |

Value

a named list of parsed options

See Also

Other internal model creation functions: [create_model.covs_and_comps\(\)](#), [create_model.formula\(\)](#), [create_model.likelihood\(\)](#)

create_plot_df	<i>Helper function for plots</i>
----------------	----------------------------------

Description

Helper function for plots

Usage

```
create_plot_df(object, x = "age", group_by = "id")
```

Arguments

object	model or fit
x	x-axis variable name
group_by	grouping variable name (use NULL for no grouping)

Value

a data frame

create_scaling	<i>Create a standardizing transform</i>
----------------	---

Description

Create a standardizing transform

Usage

```
create_scaling(x, name)
```

Arguments

x	variable measurements (might contain NA or NaN)
name	variable name

Value

an object of class [lgpscaling](#)

See Also

Other variable scaling functions: [apply_scaling\(\)](#)

dinvgamma_stanlike	<i>Density and quantile functions of the inverse gamma distribution</i>
--------------------	---

Description

Using the same parametrization as Stan. More info [here](#).

Usage

```
dinvgamma_stanlike(x, alpha, beta, log = FALSE)
```

```
qinvgamma_stanlike(p, alpha, beta)
```

Arguments

x	point where to compute the density
alpha	positive real number
beta	positive real number
log	is log-scale used?
p	quantile (must be between 0 and 1)

Value

density/quantile value

See Also

Other functions related to the inverse-gamma distribution: [plot_invgamma\(\)](#), [priors](#)

draw_pred	<i>Draw pseudo-observations from posterior or prior predictive distribution</i>
-----------	---

Description

Draw pseudo-observations from predictive distribution. If pred contains draws from the component posterior (prior) distributions, then the output is draws from the posterior (prior) predictive distribution. If pred is not specified, then whether output draws are from prior or posterior predictive distribution depends on whether fit is created using the [lgp](#) option prior_only=TRUE or not.

Usage

```
draw_pred(fit, pred = NULL)
```

Arguments

fit	An object of class lgpfit that has been created using the lgp option <code>sample_f=TRUE</code> .
pred	An object of class Prediction , containing draws of each model component. If NULL, this is obtained using <code>get_pred(fit)</code> .

Value

An array with shape $S \times P$, where S is the number of draws that `pred` contains and P is the length of each function draw. Each row $s = 1, \dots, S$ of the output is one vector drawn from the predictive distribution, given parameter draw s .

See Also

Other main functions: [create_model\(\)](#), [get_draws\(\)](#), [lgp\(\)](#), [pred\(\)](#), [prior_pred\(\)](#), [sample_model\(\)](#)

example_fit

*Quick way to create an example lgpfit, useful for debugging***Description**

Quick way to create an example `lgpfit`, useful for debugging

Usage

```
example_fit(
  formula = y ~ id + age + age | SEX + age | LOC,
  likelihood = "gaussian",
  chains = 1,
  iter = 30,
  num_indiv = 6,
  num_timepoints = 5,
  ...
)
```

Arguments

formula	model formula
likelihood	observation model
chains	number of chains to run
iter	number of iterations to run
num_indiv	number of individuals (data simulation)
num_timepoints	number of time points (data simulation)
...	additional arguments to lgp

Value

An [lgpfit](#) object created by fitting the example model.

fit_summary	<i>Print a fit summary.</i>
-------------	-----------------------------

Description

Print a fit summary.

Usage

```
fit_summary(fit, ignore_pars = c("f_latent", "eta", "teff_raw", "lp_"))
```

Arguments

fit	an object of class lgpfit
ignore_pars	parameters and generated quantities to ignore from output

Value

object invisibly.

GaussianPrediction-class

An S4 class to represent analytically computed predictive distributions (conditional on hyperparameters) of an additive GP model

Description

An S4 class to represent analytically computed predictive distributions (conditional on hyperparameters) of an additive GP model

Usage

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

```
## S4 method for signature 'GaussianPrediction'
component_names(object)
```

```
## S4 method for signature 'GaussianPrediction'
num_components(object)
```

```
## S4 method for signature 'GaussianPrediction'
num_paramsets(object)
```

```
## S4 method for signature 'GaussianPrediction'
num_evalpoints(object)
```

Arguments

object [GaussianPrediction](#) object for which to apply a class method.

Methods (by generic)

- show([GaussianPrediction](#)): Print a summary about the object.
- component_names([GaussianPrediction](#)): Get names of components.
- num_components([GaussianPrediction](#)): Get number of components.
- num_paramsets([GaussianPrediction](#)): Get number of parameter combinations (different parameter vectors) using which predictions were computed.
- num_evalpoints([GaussianPrediction](#)): Get number of points where predictions were computed.

Slots

f_comp_mean component means
 f_comp_std component standard deviations
 f_mean signal mean (on normalized scale)
 f_std signal standard deviation (on normalized scale)
 y_mean predictive mean (on original data scale)
 y_std predictive standard deviation (on original data scale)
 x a data frame of points (covariate values) where the function posteriors or predictive distributions have been evaluated

See Also

[Prediction](#)

get_draws	<i>Extract parameter draws from lgpfitt or stanfit</i>
-----------	--

Description

Uses [extract](#) with permuted = FALSE and inc_warmup = FALSE.

Usage

```
get_draws(object, draws = NULL, reduce = NULL, ...)
```

Arguments

object	An object of class lgpfitt or stanfit.
draws	Indices of the parameter draws. NULL corresponds to all post-warmup draws.
reduce	Function used to reduce all parameter draws into one set of parameters. Ignored if NULL, or if draws is not NULL.
...	Additional arguments to <code>rstan::extract()</code> .

Value

The return value is always a 2-dimensional array of shape `num_param_sets x num_params`.

See Also

Other main functions: [create_model\(\)](#), [draw_pred\(\)](#), [lgp\(\)](#), [pred\(\)](#), [prior_pred\(\)](#), [sample_model\(\)](#)

<code>get_pred</code>	<i>Extract model predictions and function posteriors</i>
-----------------------	--

Description

NOTE: It is not recommended for users to call this. Use [pred](#) instead.

Usage

```
get_pred(fit, draws = NULL, reduce = NULL, verbose = TRUE)
```

Arguments

<code>fit</code>	An object of class lgpfit .
<code>draws</code>	Indices of parameter draws to use, or <code>NULL</code> to use all draws.
<code>reduce</code>	Reduction for parameters draws. Can be a function that is applied to reduce all parameter draws into one parameter set, or <code>NULL</code> (no reduction). Has no effect if <code>draws</code> is specified.
<code>verbose</code>	Should more information and a possible progress bar be printed?

Value

an object of class [GaussianPrediction](#) or [Prediction](#)

<code>kernel</code>	<i>Compute a kernel matrix (covariance matrix)</i>
---------------------	--

Description

These have `STAN_kernel_*` counterparts. These R versions are provided for reference and are not optimized for speed. These are used when generating simulated data, and not during model inference.

Usage

```

kernel_eq(x1, x2, alpha = 1, ell)

kernel_ns(x1, x2, alpha = 1, ell, a)

kernel_zerohsum(x1, x2, M)

kernel_bin(x1, x2, pos_class = 0)

kernel_cat(x1, x2)

kernel_varmask(x1, x2, a, vm_params)

kernel_beta(beta, idx1_expand, idx2_expand)

```

Arguments

x1	vector of length n
x2	vector of length m
alpha	marginal std (default = 1)
ell	lengthscale
a	steepness of the warping function rise
M	number of categories
pos_class	binary (mask) kernel function has value one if both inputs have this value, otherwise it is zero
vm_params	vector of two mask function parameters.
beta	a parameter vector (row vector) of length N_{cases}
idx1_expand	integer vector of length n
idx2_expand	integer vector of length m

Value

A matrix of size $n \times m$.

Functions

- `kernel_eq()`: Uses the exponentiated quadratic kernel.
- `kernel_ns()`: Uses the non-stationary kernel (input warping + squared exponential).
- `kernel_zerohsum()`: Uses the zero-sum kernel. Here, x1 and x2 must be integer vectors (integers denoting different categories). Returns a binary matrix.
- `kernel_bin()`: Uses the binary (mask) kernel. Here, x1 and x2 must be integer vectors (integers denoting different categories). Returns a binary matrix.
- `kernel_cat()`: Uses the categorical kernel. Here, x1 and x2 must be integer vectors (integers denoting different categories). Returns a binary matrix.

- `kernel_varmask()`: Computes variance mask multiplier matrix. NaN's in `x1` and `x2` will be replaced by 0.
- `kernel_beta()`: Computes the heterogeneity multiplier matrix. *NOTE*: `idx_expand` needs to be given so that `idx_expand[j]-1` tells the index of the beta parameter that should be used for the j th observation. If observation j doesn't correspond to any beta parameter, then `idx_expand[j]` should be 1.

`KernelComputer-class` *An S4 class to represent input for kernel matrix computations*

Description

An S4 class to represent input for kernel matrix computations

Usage

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

## S4 method for signature 'KernelComputer'
num_components(object)

## S4 method for signature 'KernelComputer'
num_evalpoints(object)

## S4 method for signature 'KernelComputer'
num_paramsets(object)

## S4 method for signature 'KernelComputer'
component_names(object)
```

Arguments

`object` The object for which to call a class method.

Methods (by generic)

- `show(KernelComputer)`: Print a summary about the object.
- `num_components(KernelComputer)`: Get number of components.
- `num_evalpoints(KernelComputer)`: Get number of evaluation points.
- `num_paramsets(KernelComputer)`: Get number of parameter sets.
- `component_names(KernelComputer)`: Get component names.

Slots

`input` Common input (for example parameter values).

`K_input` Input for computing kernel matrices between data points ($N \times N$). A list.

`Ks_input` Input for computing kernel matrices between data and output points ($P \times N$). A list.

`Kss_input` Input for computing kernel matrices between output points ($P \times P$). A list, empty if `full_covariance=FALSE`.

`comp_names` Component names (character vector).

`full_covariance` Boolean value determining if this can compute full predictive covariance matrices (or just marginal variance at each point).

`no_separate_output_points` Boolean value determining if `Ks_input` and `Kss_input` are the same thing. Using this knowledge can reduce unnecessary computations of kernel matrices.

`STREAM` external pointer (for calling 'Stan' functions)

lgp

*Main function of the 'lgpr' package***Description**

Creates an additive Gaussian process model using `create_model` and fits it using `sample_model`. See the [Mathematical description of lgpr models](#) vignette for more information about the connection between different options and the created statistical model.

Usage

```
lgp(
  formula,
  data,
  likelihood = "gaussian",
  prior = NULL,
  c_hat = NULL,
  num_trials = NULL,
  options = NULL,
  prior_only = FALSE,
  verbose = FALSE,
  sample_f = !(likelihood == "gaussian"),
  quiet = FALSE,
  skip_postproc = sample_f,
  ...
)
```

Arguments

formula	The model formula, where • it must contain exactly one tilde (~), with response variable on the left-hand side and model terms on the right-hand side • terms are separated by a plus (+) sign • all variables appearing in formula must be found in data See the "Model formula syntax" section below (lgp) for instructions on how to specify the model terms.
data	A <code>data.frame</code> where each column corresponds to one variable, and each row is one observation. Continuous covariates and the response variable must have type "numeric" and categorical covariates must have type "factor". Missing values should be indicated with <code>NaN</code> or <code>NA</code>. The response variable cannot contain missing values. Column names should not contain trailing or leading underscores.
likelihood	Determines the observation model. Must be either "gaussian" (default), "poisson", "nb" (negative binomial), "binomial" or "bb" (beta binomial).
prior	A named list, defining the prior distribution of model (hyper)parameters. See the "Defining priors" section below (lgp).
c_hat	The GP mean. This should only be given if <code>sample_f</code> is <code>TRUE</code>, otherwise the GP will always have zero mean. If <code>sample_f</code> is <code>TRUE</code>, the given <code>c_hat</code> can be a vector of length <code>dim(data)[1]</code>, or a real number defining a constant GP mean. If not specified and <code>sample_f</code> is <code>TRUE</code>, <code>c_hat</code> is set to • <code>c_hat = mean(y)</code>, if likelihood is "gaussian", • <code>c_hat = log(mean(y))</code> if likelihood is "poisson" or "nb", • <code>c_hat = log(p/(1-p))</code>, where <code>p = mean(y/num_trials)</code> if likelihood is "binomial" or "bb", where <code>y</code> denotes the response variable measurements.
num_trials	This argument (number of trials) is only needed when likelihood is "binomial" or "bb". Must have length one or equal to the number of data points. Setting <code>num_trials=1</code> and <code>likelihood="binomial"</code> corresponds to Bernoulli observation model.
options	A named list with the following possible fields: • <code>delta</code> Amount of added jitter to ensure positive definite covariance matrices. • <code>vm_params</code> Variance mask function parameters (numeric vector of length 2). If <code>options</code> is <code>NULL</code>, default options are used. The defaults are equivalent to <code>options = list(delta = 1e-8, vm_params = c(0.025, 1))</code>.
prior_only	Should likelihood be ignored? See also sample_param_prior which can be used for any lgpmodel, and whose runtime is independent of the number of observations.
verbose	Can messages be printed during model creation? Has no effect if <code>quiet=TRUE</code>.

<code>sample_f</code>	Determines if the latent function values are sampled (must be TRUE if likelihood is not "gaussian"). If this is TRUE, the response variable will be normalized to have zero mean and unit variance.
<code>quiet</code>	Should all output messages be suppressed? You need to set also <code>refresh=0</code> if you want to suppress also the progress update messages from sampling .
<code>skip_postproc</code>	Should all postprocessing be skipped? If this is TRUE, the returned lgpfit object will likely be much smaller (if <code>sample_f=FALSE</code>).
<code>...</code>	Optional arguments passed to sampling or optimizing .

Value

Returns an object of the S4 class [lgpfit](#).

Model formula syntax

There are two ways to define the model formula:

1. Using a common [formula](#)-like syntax, like in `y ~ age + age|id + sex`. Terms can consist of a single variable, such as `age`, or an interaction of two variables, such as `age|id`. In single-variable terms, the variable can be either continuous (numeric) or categorical (factor), whereas in interaction terms the variable on the left-hand side of the vertical bar (`|`) has to be continuous and the one on the right-hand side has to be categorical. Formulae specified using this syntax are translated to the advanced format so that
 - single-variable terms become `gp(x)` if variable `x` is numeric and `zs(x)` if `x` is a factor
 - interaction terms `x|z` become `gp(x)*zs(z)`
2. Using the advanced syntax, like in `y ~ gp(age) + gp(age)*zs(id) + het(id)*gp_vm(disAge)`. This creates [lgprhs](#) objects, which consist of [lgpterms](#), which consist of [lgpexprs](#). This approach must be used if creating nonstationary, heterogeneous or temporally uncertain components.

Either one of the approaches should be used and they should not be mixed.

Defining priors

The prior argument must be a named list, like `list(alpha=student_t(4), wrp=igam(30,10))`. See examples in tutorials. Possible allowed names are

- `"alpha"` = component magnitude parameters
- `"ell"` = component lengthscale parameters
- `"wrp"` = input warping steepness parameters
- `"sigma"` = noise magnitude (Gaussian obs. model)
- `"phi"` = inv. overdispersion (negative binomial obs. model)
- `"gamma"` = overdispersion (beta-binomial obs. model)
- `"beta"` = heterogeneity parameters
- `"effect_time"` = uncertain effect time parameters
- `"effect_time_info"` = additional options for the above

See [priors](#) for functions that can be used to define the list elements. If a parameter of a model is not given in this list, a default prior will be used for it.

When to not use default priors

It is not recommended to use default priors blindly. Rather, priors should be specified according to the knowledge about the problem at hand, as in any Bayesian analysis. In `lgpr` this is especially important when

1. Using a non-Gaussian likelihood or otherwise setting `sample_f = TRUE`. In this case the response variable is not normalized, so the scale on which the data varies must be taken into account when defining priors of the signal magnitude parameters `alpha` and possible noise parameters (`sigma`, `phi`, `gamma`). Also it should be checked if `c_hat` is set in a sensible way.
2. Using a model that contains a `gp_ns(x)` or `gp_vm(x)` expression in its formula. In this case the corresponding covariate `x` is not normalized, and the prior for the input warping steepness parameter `wrp` must be set according to the expected width of the window in which the non-stationary effect of `x` occurs. By default, the width of this window is about 36, which has been set assuming that the unit of `x` is months.

See Also

Other main functions: `create_model()`, `draw_pred()`, `get_draws()`, `pred()`, `prior_pred()`, `sample_model()`

`lgpexpr-class`

An S4 class to represent an lgp expression

Description

An S4 class to represent an lgp expression

Slots

`covariate` name of a covariate

`fun` function name

See Also

See [operations](#) for performing arithmetics on `lgprhs`, `lgpterm` and `lgpexpr` objects.

lgpfit-class*An S4 class to represent the output of the lgp function*

Description

An S4 class to represent the output of the lgp function

Usage

```
## S4 method for signature 'lgpfit'  
show(object)  
  
## S4 method for signature 'lgpfit'  
component_names(object)  
  
## S4 method for signature 'lgpfit'  
num_components(object)  
  
## S4 method for signature 'lgpfit'  
postproc(object, verbose = TRUE)  
  
## S4 method for signature 'lgpfit'  
contains_postproc(object)  
  
## S4 method for signature 'lgpfit'  
clear_postproc(object)  
  
## S4 method for signature 'lgpfit'  
get_model(object)  
  
## S4 method for signature 'lgpfit'  
get_stanfit(object)  
  
## S4 method for signature 'lgpfit'  
is_f_sampled(object)  
  
## S4 method for signature 'lgpfit,missing'  
plot(x, y)
```

Arguments

object	The object for which to apply a class method.
verbose	Can the method print any messages?
x	an lgpfit object to visualize
y	unused argument

Methods (by generic)

- `show(lgpfit)`: Print information and summary about the fit object.
- `component_names(lgpfit)`: Get names of model components.
- `num_components(lgpfit)`: Get number of model components. Returns a positive integer.
- `postproc(lgpfit)`: Apply postprocessing. Returns an updated [lgpfit](#) object (copies data).
- `contains_postproc(lgpfit)`: Check if object contains postprocessing information.
- `clear_postproc(lgpfit)`: Returns an updated (copies data) [lgpfit](#) object without any post-processing information.
- `get_model(lgpfit)`: Get the stored [lgpmodel](#) object. Various properties of the returned object can be accessed as explained in the documentation of [lgpmodel](#).
- `get_stanfit(lgpfit)`: Get the stored [stanfit](#) object. Various properties of the returned object can be accessed or plotted as explained [here](#) or in the documentation of [stanfit](#).
- `is_f_sampled(lgpfit)`: Determine if inference was done by sampling the latent signal f (and its components).
- `plot(x = lgpfit, y = missing)`: Visualize parameter draws using [plot_draws](#).

Slots

`stan_fit` An object of class `stanfit`.

`model` An object of class `lgpmodel`.

`num_draws` Total number of parameter draws.

`postproc_results` A named list containing possible postprocessing results.

See Also

For extracting parameter draws, see [get_draws](#), or the `rstan` methods for `stanfit` objects.

For more detailed plotting functions, see [plot_draws](#), [plot_beta](#), [plot_warp](#), [plot_effect_times](#)

<code>lgpformula-class</code>	<i>An S4 class to represent an lgp formula</i>
-------------------------------	--

Description

An S4 class to represent an lgp formula

Slots

`terms` an object of class [lgprhs](#)

`y_name` name of the response variable

`call` original formula call

See Also

See [operations](#) for performing arithmetics on [lgprhs](#), [lgpterm](#) and [lgpexpr](#) objects.

lgpmodel-class	<i>An S4 class to represent an additive GP model</i>
----------------	--

Description

An S4 class to represent an additive GP model

Usage

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

## S4 method for signature 'lgpmodel'
parameter_info(object, digits = 3)

## S4 method for signature 'lgpmodel'
component_info(object)

## S4 method for signature 'lgpmodel'
num_components(object)

## S4 method for signature 'lgpmodel'
covariate_info(object)

## S4 method for signature 'lgpmodel'
component_names(object)

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

Arguments

object	The object for which to apply a class method.
digits	number of digits to show for floating point numbers

Methods (by generic)

- `show(lgpmodel)`: Print information and summary about the object. Returns object invisibly.
- `parameter_info(lgpmodel)`: Get a parameter summary (bounds and priors). Returns a `data.frame`.
- `component_info(lgpmodel)`: Get a data frame with information about each model component.
- `num_components(lgpmodel)`: Get number of model components. Returns a positive integer.
- `covariate_info(lgpmodel)`: Get covariate information.
- `component_names(lgpmodel)`: Get names of model components.
- `is_f_sampled(lgpmodel)`: Determine if inference of the model requires sampling the latent signal f (and its components).

Slots

formula An object of class [lgpformula](#)
data The original unmodified data.
stan_input The data to be given as input to `rstan::sampling`
var_names List of variable names grouped by type.
var_scalings A named list with fields

- **y** - Response variable normalization function and its inverse operation. Must be an [lgp-scaling](#) object.
- **x_cont** - Continuous covariate normalization functions and their inverse operations. Must be a named list with each element is an [lgpscaling](#) object.

var_info A named list with fields

- **x_cat_levels** - Names of the levels of categorical covariates before converting from factor to numeric.

info Other info in text format.
sample_f Whether the signal **f** is sampled or marginalized.
full_prior Complete prior information.

 lgprhs-class

An S4 class to represent the right-hand side of an lgp formula

Description

An S4 class to represent the right-hand side of an lgp formula

Slots

summands a list of one or more [lgpterm](#)s

See Also

See [operations](#) for performing arithmetics on [lgprhs](#), [lgpterm](#) and [lgpexpr](#) objects.

 lgpscaling-class

An S4 class to represent variable scaling

Description

An S4 class to represent variable scaling

Slots

loc original location (mean)
scale original scale (standard deviation)
var_name variable name

lgpsim-class	<i>An S4 class to represent a data set simulated using the additive GP formalism</i>
--------------	--

Description

An S4 class to represent a data set simulated using the additive GP formalism

Usage

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

## S4 method for signature 'lgpsim,missing'
plot(x, y, ...)
```

Arguments

object	an lgpsim object
x	an lgpsim object to plot
y	not used
...	optional arguments passed to plot_sim

Methods (by generic)

- `show(lgpsim)`: Show summary of object.
- `plot(x = lgpsim, y = missing)`: Plot the data and generating process. For more information see [plot_sim](#).

Slots

data	the actual data
response	name of the response variable in the data
components	the drawn function components
kernel_matrices	the covariance matrices for each gp
info	A list with fields • <code>par_ell</code> the used lengthscale parameters • <code>par_cont</code> the parameters used to generate the continuous covariates • <code>p_signal</code> signal proportion
effect_times	A list with fields • <code>true</code> possible true effect times that generate the disease effect • <code>observed</code> possible observed effect times

lgp <code>term</code> -class	<i>An S4 class to represent one formula term</i>
------------------------------	--

Description

An S4 class to represent one formula term

Slots

factors a list of at most two [lgp`exprs`](#)

See Also

See [operations](#) for performing arithmetics on [lgp`rhs`](#), [lgp`term`](#) and [lgp`expr`](#) objects.

model <code>_summary</code>	<i>Print a model summary.</i>
-----------------------------	-------------------------------

Description

Print a model summary.

Usage

```
model_summary(object, digits = 3)
```

```
param_summary(object, digits = 3)
```

Arguments

object	a model or fit
digits	number of digits to round floats to

Value

object invisibly.

new_x	<i>Create test input points for prediction</i>
-------	--

Description

Replaces a continuous variable `x` in the data frame, and possibly another continuous variable `x_ns` derived from it, with new values, for each level of a grouping factor (usually `id`)

Usage

```
new_x(data, x_values, group_by = "id", x = "age", x_ns = NULL)
```

Arguments

<code>data</code>	A data frame. Can also be an lgpfit or lgpmodel object, in which case data is extracted from it.
<code>x_values</code>	the values of <code>x</code> to set for each individual
<code>group_by</code>	name of the grouping variable, must be a factor in data (or use <code>group_by=NA</code> to create a dummy grouping factor which has only one value)
<code>x</code>	of the variable along which to extend, must be a numeric in data
<code>x_ns</code>	of a nonstationary variable derived from <code>x</code> , must be a numeric in data

Value

a data frame containing the following columns

- all factors in the original data
- `x`
- `x_ns` (unless it is `NULL`)

See Also

Other data frame handling functions: [add_dis_age\(\)](#), [add_factor\(\)](#), [add_factor_crossing\(\)](#), [adjusted_c_hat\(\)](#), [split\(\)](#)

operations

Operations on formula terms and expressions

Description

Operations on formula terms and expressions

Usage

```
## S4 method for signature 'lgprhs,lgprhs'  
e1 + e2  
  
## S4 method for signature 'lgpterm,lgpterm'  
e1 + e2  
  
## S4 method for signature 'lgprhs,lgpterm'  
e1 + e2  
  
## S4 method for signature 'lgpterm,lgpterm'  
e1 * e2
```

Arguments

e1	The first sum, term or expression
e2	The second sum, term or expression

Value

The behaviour and return type depend on the types of e1 and e2. You can

- sum (+) two [lgprhs](#)'s to yield an [lgprhs](#)
- sum (+) two [lgpterm](#)'s to yield an [lgprhs](#)
- sum (+) an [lgprhs](#) and an [lgpterm](#) to yield an [lgprhs](#)
- multiply (*) two [lgpterm](#)'s to yield an [lgpterm](#)

plot_api_c

Plot a generated/fit model component

Description

Data frames specified in arguments `df`, and `df_err` must have a format where

- The first column is the grouping factor (usually `id`).
- The second column is the x-axis variable (usually `age`).
- The third column is the coloring factor. If name of the third column is `NA`, coloring is not done.
- A column named `y` must contain the y-axis variable (not for `df_err`).
- A column named `lower` (`upper`) must contain the lower (`upper`) bound of error bar (only for `df_err`).
- The posterior draw using which the fit has been computed can be specified with a factor named `_draw_` (only for `df`).

Usage

```
plot_api_c(  
  df,  
  df_err = NULL,  
  alpha = 1,  
  alpha_err = 0.2,  
  no_err = FALSE,  
  no_line = FALSE  
)
```

Arguments

<code>df</code>	a data frame
<code>df_err</code>	a data frame
<code>alpha</code>	line opacity
<code>alpha_err</code>	ribbon opacity
<code>no_err</code>	hide error bar even when it would normally be plotted?
<code>no_line</code>	hide line even when it would normally be plotted?

Value

A `ggplot` object.

See Also

Other internal plot API functions: `plot_api_g()`

plot_api_g	<i>Plot longitudinal data and/or model fit so that each subject/group has their own panel</i>
------------	---

Description

Data frames specified in arguments `df_data`, `df_signal`, `df_fit`, and `df_fit_err` must have a format where

- the first column is the grouping factor (usually id)
- the second column is the x-axis variable (usually age)
- a column named `y` must contain the y-axis variable (not for `df_fit_err`)
- a column named `lower` (`upper`) must contain the lower (`upper`) bound of error bar (only for `df_fit_err`)
- a column named `draw` must be a factor that specifies the posterior draw using which the fit has been computed (only for `df_fit`)

Usage

```
plot_api_g(
  df_data,
  df_signal = NULL,
  df = NULL,
  df_err = NULL,
  teff_signal = NULL,
  teff_obs = NULL,
  i_test = NULL,
  color_signal = color_palette(2)[1],
  color = color_palette(2)[2],
  color_err = colorset("red", "light_highlight"),
  color_vlines = colorset("gray", "mid_highlight"),
  alpha = 1,
  alpha_err = 0.5,
  nrow = NULL,
  ncol = NULL,
  y_transform = function(x) x
)
```

Arguments

<code>df_data</code>	A data frame containing the observations.
<code>df_signal</code>	A data frame containing the true signal. Omitted if <code>NULL</code> .
<code>df</code>	A data frame containing the model fit, or a list of data frames. The list version can be used for example so that each list element corresponds to the fit computed using one parameter draw. Omitted if <code>NULL</code> .

df_err	A data frame containing error bars. Omitted if NULL. Must be NULL if df_fit is a list.
teff_signal	A named vector containing true effect times used to generate the signal. Omitted if NULL.
teff_obs	A named vector containing observed effect times. Omitted if NULL.
i_test	Indices of test points.
color_signal	Line color for true signal.
color	Line color for model fit.
color_err	Color of the error ribbon.
color_vlines	Two line colors for vertical lines (true and obs. effect time).
alpha	Line opacity for model fit.
alpha_err	Opacity of the error ribbon.
nrow	number of rows, an argument for facet_wrap
ncol	number of columns, an argument for facet_wrap
y_transform	A function to be applied to the third column of df_data.

Value

A [ggplot](#) object.

See Also

Other internal plot API functions: [plot_api_c\(\)](#)

plot_components	<i>Visualize all model components</i>
-----------------	---------------------------------------

Description

This calls [plot_f](#) for all model components.

Usage

```
plot_components(
  fit,
  pred = NULL,
  group_by = "id",
  t_name = "age",
  MULT_STD = 2,
  verbose = TRUE,
  draws = NULL,
  reduce = function(x) base::mean(x),
  color_by = NA,
  no_err = FALSE,
```

```

ylim = NULL,
draw = TRUE,
nrow = NULL,
ncol = NULL,
gg_add = NULL,
x = NULL,
...
)

```

Arguments

fit	An object of class lgpfit .
pred	An object of class GaussianPrediction or Prediction . If pred=NULL, the pred function is called with the given reduce and draws arguments.
group_by	name of the grouping variable (use group_by=NA to avoid grouping)
t_name	name of the x-axis variable
MULT_STD	a multiplier for standard deviation
verbose	Can this print any messages?
draws	Only has effect if pred=NULL.
reduce	Only has effect if pred=NULL.
color_by	Names of coloring factors. Can have length 1 or equal to the number of components. See the color_by argument of plot_f .
no_err	Should the error ribbons be skipped even though they otherwise would be shown? Can have length 1 or equal to number of components + 1. See the no_err argument of plot_api_c .
ylim	a vector of length 2 (upper and lower y-axis limits), or NULL
draw	if this is TRUE, the plot grid is drawn using arrangeGrob
nrow	number of grid rows
ncol	number of grid columns
gg_add	additional ggplot object to add to each plot
x	Deprecated argument. This is now taken from the pred object to ensure compatibility.
...	additional arguments to plot_api_c

Value

a list of ggplot objects invisibly

See Also

Other main plot functions: [plot_draws\(\)](#), [plot_pred\(\)](#)

`plot_data`*Vizualizing longitudinal data*

Description

Vizualizing longitudinal data

Usage

```
plot_data(  
  data,  
  x_name = "age",  
  y_name = "y",  
  group_by = "id",  
  facet_by = NULL,  
  color_by = NULL,  
  highlight = NULL,  
  main = NULL,  
  sub = NULL  
)
```

Arguments

<code>data</code>	A data frame.
<code>x_name</code>	Name of x-axis variable.
<code>y_name</code>	Name of the y-axis variable.
<code>group_by</code>	Name of grouping variable (must be a factor).
<code>facet_by</code>	Name of the faceting variable (must be a factor).
<code>color_by</code>	Name of coloring variable (must be a factor).
<code>highlight</code>	Value of category of the <code>group_by</code> variable that is highlighted. Can only be used if <code>color_by</code> is NULL.
<code>main</code>	main plot title
<code>sub</code>	plot subtitle

Value

a `ggplot` object

plot_draws

*Visualize the distribution of parameter draws***Description**

Visualize the distribution of parameter draws

Usage

```
plot_draws(
  fit,
  type = "intervals",
  regex_pars = c("alpha", "ell", "wrp", "sigma", "phi", "gamma"),
  ...
)
```

```
plot_beta(fit, type = "dens", verbose = TRUE, ...)
```

```
plot_warp(
  fit,
  num_points = 300,
  window_size = 48,
  color = colorset("red", "dark"),
  alpha = 0.5
)
```

```
plot_effect_times(fit, type = "areas", verbose = TRUE, ...)
```

Arguments

fit	an object of class lgpfit
type	plot type, allowed options are "intervals", "dens", "areas", and "trace"
regex_pars	regex for parameter names to plot
...	additional arguments for the bayesplot function mcmc_intervals , mcmc_dens , mcmc_areas or mcmc_trace
verbose	Can any output be printed?
num_points	number of plot points
window_size	width of time window
color	line color
alpha	line alpha

Value

a ggplot object or list of them

Functions

- `plot_draws()`: visualizes the distribution of any set of model parameters (defaults to kernel hyperparameters and possible observation model parameters)
- `plot_beta()`: visualizes the distribution of the individual-specific disease effect magnitude parameter draws
- `plot_warp()`: visualizes the input warping function for different draws of the warping steepness parameter
- `plot_effect_times()`: visualizes the input warping function for different parameter draws

See Also

Other main plot functions: [plot_components\(\)](#), [plot_pred\(\)](#)

<code>plot_inputwarp</code>	<i>Visualize input warping function with several steepness parameter values</i>
-----------------------------	---

Description

Visualize input warping function with several steepness parameter values

Usage

```
plot_inputwarp(wrp, x, color = colorset("red", "dark"), alpha = 0.5)
```

Arguments

<code>wrp</code>	a vector of values of the warping steepness parameter
<code>x</code>	a vector of input values
<code>color</code>	line color
<code>alpha</code>	line alpha

Value

a ggplot object

plot_invgamma

Plot the inverse gamma-distribution pdf

Description

Plot the inverse gamma-distribution pdf

Usage

```
plot_invgamma(  
  alpha,  
  beta,  
  by = 0.01,  
  log = FALSE,  
  IQR = 0.95,  
  return_quantiles = FALSE,  
  linecolor = colorset("red", "dark"),  
  fillcolor = colorset("red", "mid")  
)
```

Arguments

alpha	positive real number
beta	positive real number
by	grid size
log	is log-scale used?
IQR	inter-quantile range width
return_quantiles	should this return a list
linecolor	line color
fillcolor	fill color

Value

a ggplot object

See Also

Other functions related to the inverse-gamma distribution: [dinvgamma_stanlike\(\)](#), [priors](#)

plot_pred

*Visualizing model predictions or inferred covariate effects***Description**

- Function draws at data points can be visualized using plot_pred. If the pred argument is NULL, it is computed using the [pred](#) function with x=NULL.
- The total signal f or any of its additive components can be plotted using plot_f.

Usage

```
plot_pred(
  fit,
  pred = NULL,
  group_by = "id",
  t_name = "age",
  MULT_STD = 2,
  verbose = TRUE,
  draws = NULL,
  reduce = function(x) base::mean(x),
  x = NULL,
  ...
)
```

```
plot_f(
  fit,
  pred = NULL,
  group_by = "id",
  t_name = "age",
  MULT_STD = 2,
  verbose = TRUE,
  draws = NULL,
  reduce = function(x) base::mean(x),
  comp_idx = NULL,
  color_by = NA,
  x = NULL,
  ...
)
```

Arguments

fit	An object of class lgpfit .
pred	An object of class GaussianPrediction or Prediction . If pred=NULL, the pred function is called with the given reduce and draws arguments.
group_by	name of the grouping variable (use group_by=NA to avoid grouping)
t_name	name of the x-axis variable

MULT_STD	a multiplier for standard deviation
verbose	Can this print any messages?
draws	Only has effect if pred=NULL.
reduce	Only has effect if pred=NULL.
x	Deprecated argument. This is now taken from the pred object to ensure compatibility.
...	additional arguments to plot_api_g or plot_api_c
comp_idx	Index of component to plot. The total sum is plotted if this is NULL.
color_by	name of coloring factor

Value

a [ggplot](#) object

See Also

Other main plot functions: [plot_components\(\)](#), [plot_draws\(\)](#)

plot_sim

Visualize an lgpsim object (simulated data)

Description

Visualize an lgpsim object (simulated data)

Usage

```
plot_sim(
  simdata,
  group_by = "id",
  x_name = "age",
  h_name = "h",
  y_name = "y",
  comp_idx = NULL,
  color_by = NA,
  verbose = TRUE,
  ...
)
```

Arguments

<code>simdata</code>	an object of class lgpsim
<code>group_by</code>	grouping factor
<code>x_name</code>	name of x-axis variable
<code>h_name</code>	name of the signal in <code>simdata\$components</code> ("h" or "f")
<code>y_name</code>	name of response variable
<code>comp_idx</code>	Possible index of a component to be shown. If this is NULL, the data and total signal are shown.
<code>color_by</code>	coloring factor
<code>verbose</code>	should some information be printed?
<code>...</code>	additional arguments to plot_api_g or plot_api_c

Value

a [ggplot](#) object

ppc

Graphical posterior predictive checks

Description

Graphical posterior predictive checks

Usage

```
ppc(fit, data = NULL, fun = default_ppc_fun(fit), verbose = TRUE, ...)
```

Arguments

<code>fit</code>	An object of class lgpfit that can be created with <code>sample_f=TRUE</code> .
<code>data</code>	the original data frame (deprecated argument with no effect, now obtained from fit object)
<code>fun</code>	bayesplot function name
<code>verbose</code>	Can this print any messages?
<code>...</code>	additional arguments passed to the default pp_check method in bayesplot

Value

a [ggplot](#) object

See Also

Introduction to graphical posterior predictive checks: [here](#). Prior predictive check can be done by calling [prior_pred](#) and then `bayesplot::pp_check()`.

pred

*Posterior predictions and function posteriors***Description**

- If `fit` is for a model that marginalizes the latent signal f (i.e. `is_f_sampled(fit)` is `FALSE`), this computes the analytic conditional posterior distributions of each model component, their sum, and the conditional predictive distribution. All these are computed for each (hyper)parameter draw (defined by `draws`), or other parameter set (obtained by a reduction defined by `reduce`). Results are stored in a [GaussianPrediction](#) object which is then returned.
- If `fit` is for a model that samples the latent signal f (i.e. `is_f_sampled(fit)` is `TRUE`), this will extract these function samples, compute their sum, and a version of the sum f that is transformed through the inverse link function. If `x` is not `NULL`, the function draws are extrapolated to the points specified by `x` using kernel regression. Results are stored in a [Prediction](#) object which is then returned.

Usage

```
pred(
  fit,
  x = NULL,
  reduce = function(x) base::mean(x),
  draws = NULL,
  verbose = TRUE,
  STREAM = get_stream(),
  c_hat_pred = NULL,
  force = FALSE,
  debug_kc = FALSE
)
```

Arguments

<code>fit</code>	An object of class lgpfit .
<code>x</code>	A data frame of points where function posterior distributions and predictions should be computed or sampled. The function new_x provides an easy way to create it. If this is <code>NULL</code> , the data points are used.
<code>reduce</code>	Reduction for parameters draws. Can be a function that is applied to reduce all parameter draws into one parameter set, or <code>NULL</code> (no reduction). Has no effect if <code>draws</code> is specified.
<code>draws</code>	Indices of parameter draws to use, or <code>NULL</code> to use all draws.
<code>verbose</code>	Should more information and a possible progress bar be printed?
<code>STREAM</code>	External pointer. By default obtained with <code>rstan::get_stream()</code> .
<code>c_hat_pred</code>	This is only used if the latent signal f was sampled. This input contains the values added to the sum f before passing through inverse link function. Must be a vector with length equal to the number of prediction points. If original

	c_hat was constant, then c_hat_pred can be ignored, in which case this will by default use the same constant.
force	This is by default FALSE to prevent unintended large computations that might crash R or take forever. Set it to TRUE try computing no matter what.
debug_kc	If this is TRUE, this only returns a KernelComputer object that is created internally. Meant for debugging.

Value

An object of class [GaussianPrediction](#) or [Prediction](#).

See Also

Other main functions: [create_model\(\)](#), [draw_pred\(\)](#), [get_draws\(\)](#), [lgp\(\)](#), [prior_pred\(\)](#), [sample_model\(\)](#)

Prediction-class	<i>An S4 class to represent prior or posterior draws from an additive function distribution.</i>
------------------	--

Description

An S4 class to represent prior or posterior draws from an additive function distribution.

Usage

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

## S4 method for signature 'Prediction'
component_names(object)

## S4 method for signature 'Prediction'
num_components(object)

## S4 method for signature 'Prediction'
num_paramsets(object)

## S4 method for signature 'Prediction'
num_evalpoints(object)
```

Arguments

object [Prediction](#) object for which to apply a class method.

Methods (by generic)

- `show(Prediction)`: Print a summary about the object.
- `component_names(Prediction)`: Get names of components.
- `num_components(Prediction)`: Get number of components.
- `num_paramsets(Prediction)`: Get number of parameter combinations (different parameter vectors) using which predictions were computed.
- `num_evalpoints(Prediction)`: Get number of points where predictions were computed.

Slots

`f_comp` component draws

`f` signal draws

`h` predictions (signal draws + scaling factor `c_hat`, transformed through inverse link function)

`x` a data frame of points (covariate values) where the functions/predictions have been evaluated/sampled
`extrapolated` Boolean value telling if the function draws are original MCMC draws or if they have been created by extrapolating such draws.

See Also

[GaussianPrediction](#)

priors

Prior definitions

Description

These use the same parametrizations as defined in the 'Stan' documentation. See the docs for [gamma](#) and [inverse gamma](#) distributions.

Usage

`uniform(square = FALSE)`

`normal(mu, sigma, square = FALSE)`

`student_t(nu, square = FALSE)`

`gam(shape, inv_scale, square = FALSE)`

`igam(shape, scale, square = FALSE)`

`log_normal(mu, sigma, square = FALSE)`

`bet(a, b)`

Arguments

square	is prior for a square-transformed parameter?
mu	mean
sigma	standard deviation
nu	degrees of freedom
shape	shape parameter (alpha)
inv_scale	inverse scale parameter (beta)
scale	scale parameter (beta)
a	shape parameter
b	shape parameter

Value

a named list

See Also

Other functions related to the inverse-gamma distribution: [dinvgamma_stanlike\(\)](#), [plot_invgamma\(\)](#)

Examples

```
# Log-normal prior
log_normal(mu = 1, sigma = 1)

# Cauchy prior
student_t(nu = 1)

# Exponential prior with rate = 0.1
gam(shape = 1, inv_scale = 0.1)

# Create a similar priors as in LongP (Cheng et al., 2019)
# Not recommended, because a lengthscale close to 0 is possible.
a <- log(1) - log(0.1)
log_normal(mu = 0, sigma = a / 2) # for continuous lengthscale
student_t(nu = 4) # for interaction lengthscale
igam(shape = 0.5, scale = 0.005, square = TRUE) # for sigma
```

prior_pred	<i>Prior (predictive) sampling</i>
------------	------------------------------------

Description

These functions take an [lgpmodel](#) object, and

- `prior_pred` samples from the prior predictive distribution of the model
- `sample_param_prior` samples only its parameter prior using [sampling](#)

Usage

```
prior_pred(
  model,
  verbose = TRUE,
  quiet = FALSE,
  refresh = 0,
  STREAM = get_stream(),
  ...
)

sample_param_prior(model, verbose = TRUE, quiet = FALSE, ...)
```

Arguments

model	An object of class lgpmodel .
verbose	Should more information and a possible progress bar be printed?
quiet	This forces verbose to be FALSE. If you want to suppress also the output from Stan, give the additional argument refresh=0.
refresh	Argument for sampling .
STREAM	External pointer. By default obtained with <code>rstan::get_stream()</code> .
...	Additional arguments for sampling .

Value

- `prior_pred` returns a list with components
 - `y_draws`: A matrix containing the prior predictive draws as rows. Can be passed to `bayesplot::pp_check()` for graphical prior predictive checking.
 - `pred_draws`: an object of class [Prediction](#), containing prior draws of each model component and their sum
 - `param_draws`: a `stanfit` object of prior parameter draws (obtained by calling `sample_param_prior` internally)
- `sample_param_prior` returns an object of class [stanfit](#)

See Also

Other main functions: [create_model\(\)](#), [draw_pred\(\)](#), [get_draws\(\)](#), [lgp\(\)](#), [pred\(\)](#), [sample_model\(\)](#)

prior_to_num

Convert given prior to numeric format

Description

Convert given prior to numeric format

Usage

```
prior_to_num(desc)
```

Arguments

desc Prior description as a named list, containing fields

- dist - Distribution name. Must be one of 'uniform', 'normal', 'student-t', 'gamma', 'inv-gamma', or 'log-normal' (case-insensitive)
- square - Is the prior for a square-transformed parameter.

Other list fields are interpreted as hyperparameters.

Value

a named list of parsed options

read_proteomics_data *Function for reading the built-in proteomics data*

Description

Function for reading the built-in proteomics data

Usage

```
read_proteomics_data(parentDir = NULL, protein = NULL, verbose = TRUE)
```

Arguments

parentDir Path to local parent directory for the data. If this is NULL, data is downloaded from <https://github.com/jtimonen/lgpr-usage/tree/master/data/proteomics>.

protein Index or name of protein.

verbose Can this print some output?

Value

a data.frame

relevances	<i>Assess component relevances</i>
------------	------------------------------------

Description

Assess component relevances

Usage

```
relevances(fit, reduce = function(x) base::mean(x), verbose = TRUE, ...)
```

Arguments

fit	an object of class <code>lgpfit</code>
reduce	a function to apply to reduce the relevances given each parameter draw into one value
verbose	Can this print any messages?
...	currently has no effect

Value

a named vector with length equal to `num_comps + 1`

s4_generics	<i>S4 generics for <code>lgpfit</code>, <code>lgpmodel</code>, and other objects</i>
-------------	--

Description

S4 generics for `lgpfit`, `lgpmodel`, and other objects

Usage

```
parameter_info(object, digits)

component_info(object)

covariate_info(object)

component_names(object)

get_model(object)

is_f_sampled(object)

get_stanfit(object)
```

```

postproc(object, ...)

contains_postproc(object)

clear_postproc(object)

num_paramsets(object)

num_evalpoints(object)

num_components(object)

```

Arguments

<code>object</code>	object for which to apply the generic
<code>digits</code>	number of digits to show
<code>...</code>	additional optional arguments to pass

Value

- `parameter_info` returns a data frame with one row for each parameter and columns for parameter name, parameter bounds, and the assigned prior
- `component_info` returns a data frame with one row for each model component, and columns encoding information about model components
- `covariate_info` returns a list with names `continuous` and `categorical`, with information about both continuous and categorical covariates
- `component_names` returns a character vector with component names
- `get_model` for [lgpfit](#) objects returns an [lgpmodel](#)
- `is_f_sampled` returns a logical value
- `get_stanfit` returns a stanfit (rstan)
- `postproc` applies postprocessing and returns an updated [lgpfit](#)
- `clear_postproc` removes postprocessing information and returns an updated [lgpfit](#)
- `num_paramsets`, `num_evalpoints` and `num_components` return an integer

Functions

- `parameter_info()`: Get parameter information (priors etc.).
- `component_info()`: Get component information.
- `covariate_info()`: Get covariate information.
- `component_names()`: Get component names.
- `get_model()`: Get [lgpmodel](#) object.
- `is_f_sampled()`: Determine if signal `f` is sampled or marginalized.
- `get_stanfit()`: Extract stanfit object.

- `postproc()`: Perform postprocessing.
- `contains_postproc()`: Determine if object contains postprocessing information.
- `clear_postproc()`: Clear postprocessing information (to reduce size of object).
- `num_paramsets()`: Get number of parameter sets.
- `num_evalpoints()`: Get number of points where posterior is evaluated.
- `num_components()`: Get number of model components.

See Also

To find out which methods have been implemented for which classes, see [lgpfit](#), [lgpmodel](#), [Prediction](#) and [GaussianPrediction](#).

sample_model	<i>Fitting a model</i>
--------------	------------------------

Description

- `sample_model` takes an [lgpmodel](#) object and fits it using [sampling](#).
- `optimize_model` takes an [lgpmodel](#) object and fits it using [optimizing](#).

Usage

```
sample_model(
  model,
  verbose = TRUE,
  quiet = FALSE,
  skip_postproc = is_f_sampled(model),
  ...
)

optimize_model(model, ...)
```

Arguments

<code>model</code>	An object of class lgpmodel .
<code>verbose</code>	Can messages be printed?
<code>quiet</code>	Should all output messages be suppressed? You need to set also <code>refresh=0</code> if you want to suppress also the progress update messages from sampling .
<code>skip_postproc</code>	Should all postprocessing be skipped? If this is TRUE, the returned lgpfit object will likely be much smaller (if <code>sample_f=FALSE</code>).
<code>...</code>	Optional arguments passed to sampling or optimizing .

Value

- `sample_model` returns an object of class `lgpfit` containing the parameter draws, the original model object, and possible postprocessing results. See documentation of `lgpfit` for more information.
- `optimize_model` directly returns the list returned by `optimizing`. See its documentation for more information.

See Also

Other main functions: `create_model()`, `draw_pred()`, `get_draws()`, `lgp()`, `pred()`, `prior_pred()`

select	<i>Select relevant components</i>
--------	-----------------------------------

Description

- `select` performs strict selection, returning either TRUE or FALSE for each component.
- `select.integrate` is like `select`, but instead of a fixed threshold, computes probabilistic selection by integrating over a threshold density.
- `select_freq` performs the selection separately using each parameter draw and returns the frequency at which each component was selected.
- `select_freq.integrate` is like `select_freq`, but instead of a fixed threshold, computes probabilistic selection frequencies by integrating over a threshold density.

Usage

```
select(fit, reduce = function(x) base::mean(x), threshold = 0.95, ...)
```

```
select_freq(fit, threshold = 0.95, ...)
```

```
select.integrate(
  fit,
  reduce = function(x) base::mean(x),
  p = function(x) stats::dbeta(x, 100, 5),
  h = 0.01,
  verbose = TRUE,
  ...
)
```

```
select_freq.integrate(
  fit,
  p = function(x) stats::dbeta(x, 100, 5),
  h = 0.01,
  verbose = TRUE,
  ...
)
```

Arguments

<code>fit</code>	An object of class <code>lgpfit</code> .
<code>reduce</code>	The reduce argument for relevances .
<code>threshold</code>	Threshold for relevance sum. Must be a value between 0 and 1.
<code>...</code>	Additional arguments to relevances .
<code>p</code>	A threshold density over interval [0,1].
<code>h</code>	A discretization parameter for computing a quadrature.
<code>verbose</code>	Should this show a progress bar?

Value

See description.

<code>show</code>	<i>Printing formula object info using the show generic</i>
-------------------	--

Description

Printing formula object info using the show generic

Usage

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

## S4 method for signature 'lgprhs'
show(object)

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

Arguments

<code>object</code>	an object of some S4 class
---------------------	----------------------------

Value

the object invisibly

sim.create_f

*Simulate latent function components for longitudinal data analysis***Description**

Simulate latent function components for longitudinal data analysis

Usage

```
sim.create_f(
  X,
  covariates,
  relevances,
  lengthscales,
  X_affected,
  dis_fun,
  bin_kernel,
  steepness,
  vm_params,
  force_zeromean
)
```

Arguments

X	input data matrix (generated by sim.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 <code>length(relevances) = 2 + length(covariates)</code> . First two values define the relevance of the individual-specific age and shared age component, respectively.
lengthscales	A vector so that <code>length(lengthscales) = 2 + sum(covariates %in% c(0,1,2))</code> .
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 <code>dis_fun</code> 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 component 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 component

sim.create_x	<i>Create an input data frame X for simulated data</i>
--------------	--

Description

Create an input data frame X for simulated data

Usage

```
sim.create_x(
  N,
  covariates,
  names,
  n_cats,
  t_data,
  t_jitter,
  t_effect_range,
  continuous_info
)
```

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_cats	An integer vector defining the number of categories for each categorical covariate, so that <code>length(n_cats)</code> equals to the number of 2's and 3's in the covariates vector.
t_data	Measurement times (same for each individual, unless <code>t_jitter > 0</code> in which case they are perturbed).
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. Alternatively, 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 <code>lambda</code> and <code>mu</code> , which have length 3. The continuous covariates are generated so that $x \leftarrow \sin(a \cdot t + b) + c$, where <code>t <- seq(0, 2*pi, length.out = k)</code> <code>a <- mu[1] + lambda[1]*stats::runif(1)</code> <code>b <- mu[2] + lambda[2]*stats::runif(1)</code> <code>c <- mu[3] + lambda[3]*stats::runif(1)</code>

Value

a list

sim.create_y	<i>Simulate noisy observations</i>
--------------	------------------------------------

Description

Simulate noisy observations

Usage

```
sim.create_y(noise_type, f, snr, phi, gamma, N_trials)
```

Arguments

noise_type	Either "gaussian", "poisson", "nb" (negative binomial), "binomial", or "bb" (beta-binomial).
f	The underlying signal.
snr	The desired signal-to-noise ratio. This argument is valid only when <code>noise_type</code> is "gaussian".
phi	The inverse overdispersion parameter for negative binomial data. The variance is $g + g^2/\phi$.
gamma	The dispersion parameter for beta-binomial data.
N_trials	The number of trials parameter for binomial data.

Value

A list out, where

- out\$h is f mapped through an inverse link function (times N_trials if noise_type is binomial or beta-binomial)
- out\$y is the noisy response variable.

sim.kernels

Compute all kernel matrices when simulating data

Description

Compute all kernel matrices when simulating data

Usage

```
sim.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
X_affected	which individuals are affected by the disease
bin_kernel	whether or not binary (mask) kernel should be used for categorical covariates (if not, the zerosum kernel is used)
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

simulate_data	<i>Generate an artificial longitudinal data set</i>
---------------	---

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,
  gamma = 0.2,
  N_affected = round(N/2),
  t_effect_range = "auto",
  t_observed = "after_0",
  c_hat = 0,
  dis_fun = "gp_warp_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,
  force_zeromean = TRUE
)
```

Arguments

N	Number of individuals.
t_data	Measurement times (same for each individual, unless t_jitter > 0 in which case they are perturbed).
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 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 $\text{length}(\text{relevances}) = 2 + \text{length}(\text{covariates})$. First two values define the relevance of the individual-specific age and shared age component, respectively.
n_cats	An integer vector defining the number of categories for each categorical covariate, so that $\text{length}(\text{n_cats})$ 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 $\text{length}(\text{lengthscales}) = 2 + \text{sum}(\text{covariates} \%in\% c(0,1,2))$.
f_var	variance of f
noise_type	Either "gaussian", "poisson", "nb" (negative binomial), "binomial", or "bb" (beta-binomial).
snr	The desired signal-to-noise ratio. This argument is valid only when noise_type is "gaussian".
phi	The inverse overdispersion parameter for negative binomial data. The variance is $g + g^2/\phi$.
gamma	The dispersion parameter for beta-binomial data.
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. Alternatively, 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 component 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 $x \leftarrow \sin(a \cdot t + b) + c$, where • $t \leftarrow \text{seq}(0, 2 \cdot \pi, \text{length.out} = k)$ • $a \leftarrow \mu[1] + \lambda[1] \cdot \text{stats::runif}(1)$ • $b \leftarrow \mu[2] + \lambda[2] \cdot \text{stats::runif}(1)$ • $c \leftarrow \mu[3] + \lambda[3] \cdot \text{stats::runif}(1)$
N_trials	The number of trials parameter for binomial data.
force_zeromean	Should each component (excluding the disease age component) be forced to have a zero mean?

Value

An object of class [lgpsim](#).

Examples

```
# Generate Gaussian data
dat <- simulate_data(N = 4, t_data = c(6, 12, 24, 36, 48), snr = 3)

# Generate negative binomially (NB) distributed count data
dat <- simulate_data(
  N = 6, t_data = seq(2, 10, by = 2), noise_type = "nb",
  phi = 2
)
```

split

Split data into training and test sets

Description

- `split_by_factor` splits according to given factor
- `split_within_factor` splits according to given data point indices within the same level of a factor
- `split_within_factor_random` selects k points from each level of a factor uniformly at random as test data
- `split_random` splits uniformly at random
- `split_data` splits according to given data rows

Usage

```

split_by_factor(data, test, var_name = "id")

split_within_factor(data, idx_test, var_name = "id")

split_within_factor_random(data, k_test = 1, var_name = "id")

split_random(data, p_test = 0.2, n_test = NULL)

split_data(data, i_test, sort_ids = TRUE)

```

Arguments

<code>data</code>	a data frame
<code>test</code>	the levels of the factor that will be used as test data
<code>var_name</code>	name of a factor in the data
<code>idx_test</code>	indices point indices with the factor
<code>k_test</code>	desired number of test data points per each level of the factor
<code>p_test</code>	desired proportion of test data
<code>n_test</code>	desired number of test data points (if NULL, <code>p_test</code> is used to compute this)
<code>i_test</code>	test data row indices
<code>sort_ids</code>	should the test indices be sorted into increasing order

Value

a named list with names `train`, `test`, `i_train` and `i_test`

See Also

Other data frame handling functions: [add_dis_age\(\)](#), [add_factor\(\)](#), [add_factor_crossing\(\)](#), [adjusted_c_hat\(\)](#), [new_x\(\)](#)

testdata_001

A very small artificial test data, used mostly for unit tests

Description

A very small artificial test data, used mostly for unit tests

Usage

```
testdata_001
```

Format

A data frame with 24 rows and 6 variables:

id individual id, a factor with levels: 1, 2, 3, 4

age age

dis_age disease-related age

blood a continuous variable

sex a factor with 2 levels: Male, Female

y a continuous variable

See Also

Other built-in datasets: [testdata_002](#)

testdata_002

Medium-size artificial test data, used mostly for tutorials

Description

Medium-size artificial test data, used mostly for tutorials

Usage

```
testdata_002
```

Format

A data frame with 96 rows and 6 variables:

id individual id, a factor with levels: 01-12

age age

diseaseAge disease-related age

sex a factor with 2 levels: Male, Female

group a factor with 2 levels: Case, Control

y a continuous variable

See Also

[read_proteomics_data](#)

Other built-in datasets: [testdata_001](#)

validate	<i>Validate S4 class objects</i>
----------	----------------------------------

Description

Validate S4 class objects

Usage

```
validate_lgpexpr(object)
validate_lgpformula(object)
validate_lgpscaling(object)
validate_lgpfit(object)
validate_GaussianPrediction(object)
validate_Prediction(object)
```

Arguments

object an object to validate

Value

TRUE if valid, otherwise reasons for invalidity

var_mask	<i>Variance masking function</i>
----------	----------------------------------

Description

Variance masking function

Usage

```
var_mask(x, stp)
```

Arguments

x a vector of length n
stp a positive real number (steepness of mask function)

Value

a vector of length n

See Also

Other kernel utility functions: [warp_input\(\)](#)

warp_input

Input warping function

Description

Input warping function

Usage

```
warp_input(x, a)
```

Arguments

x	a vector of length n
a	steepness of the warping function rise

Value

a vector of warped inputs $w(x)$, length n

See Also

Other kernel utility functions: [var_mask\(\)](#)

Index

- * **Bayesian**
 - lgpr-package, 3
- * **GP**
 - lgpr-package, 3
- * **Stan**
 - lgpr-package, 3
- * **additive**
 - lgpr-package, 3
- * **built-in datasets**
 - testdata_001, 64
 - testdata_002, 65
- * **built-in data**
 - read_proteomics_data, 51
- * **covariate**
 - lgpr-package, 3
- * **data frame handling functions**
 - add_dis_age, 5
 - add_factor, 6
 - add_factor_crossing, 6
 - adjusted_c_hat, 7
 - new_x, 33
 - split, 63
- * **data frame handling**
 - plot_data, 39
- * **datasets**
 - testdata_001, 64
 - testdata_002, 65
- * **functions related to the inverse-gamma distribution**
 - dinvgamma_stanlike, 16
 - plot_invgamma, 42
 - priors, 48
- * **internal model creation functions**
 - create_model.covs_and_comps, 10
 - create_model.formula, 11
 - create_model.likelihood, 12
 - create_model.prior, 14
- * **internal plot API functions**
 - plot_api_c, 34
 - plot_api_g, 36
- * **interpretable**
 - lgpr-package, 3
- * **kernel utility functions**
 - var_mask, 66
 - warp_input, 67
- * **longitudinal**
 - lgpr-package, 3
- * **main functions**
 - create_model, 9
 - draw_pred, 16
 - get_draws, 19
 - lgp, 23
 - pred, 46
 - prior_pred, 49
 - sample_model, 54
- * **main plot functions**
 - plot_components, 37
 - plot_draws, 40
 - plot_pred, 43
- * **model**
 - lgpr-package, 3
- * **relevance**
 - lgpr-package, 3
- * **variable scaling functions**
 - apply_scaling, 8
 - create_scaling, 15
- *, lgpterm, lgpterm-method (operations), 34
- +, lgprhs, lgprhs-method (operations), 34
- +, lgprhs, lgpterm-method (operations), 34
- +, lgpterm, lgpterm-method (operations), 34
- add_dis_age, 4, 5, 6, 7, 33, 64
- add_factor, 4, 5, 6, 7, 33, 64
- add_factor_crossing, 4-6, 6, 7, 33, 64
- adjusted_c_hat, 4-7, 7, 33, 64
- apply_scaling, 8, 15
- arrangeGrob, 38

- as.character, lgpexpr-method
(as_character), 8
- as.character, lgpformula-method
(as_character), 8
- as.character, lgpterm-method
(as_character), 8
- as_character, 8
- bet (priors), 48
- clear_postproc (s4_generics), 52
- clear_postproc, lgpfit-method
(lgpfit-class), 27
- component_info (s4_generics), 52
- component_info, lgpmodel-method
(lgpmodel-class), 29
- component_names (s4_generics), 52
- component_names, GaussianPrediction-method
(GaussianPrediction-class), 18
- component_names, KernelComputer-method
(KernelComputer-class), 22
- component_names, lgpfit-method
(lgpfit-class), 27
- component_names, lgpmodel-method
(lgpmodel-class), 29
- component_names, Prediction-method
(Prediction-class), 47
- contains_postproc (s4_generics), 52
- contains_postproc, lgpfit-method
(lgpfit-class), 27
- covariate_info (s4_generics), 52
- covariate_info, lgpmodel-method
(lgpmodel-class), 29
- create_model, 4, 9, 17, 20, 23, 26, 47, 50, 55
- create_model.covs_and_comps, 10, 12–14
- create_model.formula, 11, 11, 13, 14
- create_model.likelihood, 11, 12, 12, 14
- create_model.options, 13
- create_model.prior, 11–13, 14
- create_plot_df, 15
- create_scaling, 8, 15
- dinvgamma_stanlike, 16, 42, 49
- draw_pred, 10, 16, 20, 26, 47, 50, 55
- example_fit, 17
- extract, 19
- facet_wrap, 37
- fit_summary, 18
- formula, 25
- gam (priors), 48
- gam, (priors), 48
- GaussianPrediction, 19, 20, 38, 43, 46–48, 54
- GaussianPrediction
(GaussianPrediction-class), 18
- GaussianPrediction-class, 18
- get_draws, 10, 17, 19, 26, 28, 47, 50, 55
- get_model (s4_generics), 52
- get_model, lgpfit-method (lgpfit-class), 27
- get_pred, 20
- get_stanfit (s4_generics), 52
- get_stanfit, lgpfit-method
(lgpfit-class), 27
- ggplot, 35, 37, 39, 44, 45
- igam (priors), 48
- igam, (priors), 48
- is_f_sampled (s4_generics), 52
- is_f_sampled, lgpfit-method
(lgpfit-class), 27
- is_f_sampled, lgpmodel-method
(lgpmodel-class), 29
- kernel, 20
- kernel_beta (kernel), 20
- kernel_bin (kernel), 20
- kernel_cat (kernel), 20
- kernel_eq (kernel), 20
- kernel_ns (kernel), 20
- kernel_varmask (kernel), 20
- kernel_zerohsum (kernel), 20
- KernelComputer, 47
- KernelComputer (KernelComputer-class), 22
- KernelComputer-class, 22
- lgp, 4, 9, 10, 12, 14, 16, 17, 20, 23, 24, 47, 50, 55
- lgpexpr, 25, 26, 28, 30, 32
- lgpexpr (lgpexpr-class), 26
- lgpexpr-class, 26
- lgpfit, 4, 17–20, 25, 27, 28, 33, 38, 40, 43, 45, 46, 53–55
- lgpfit (lgpfit-class), 27

- lgpfit-class, 27
- lgpformula, 11, 12, 30
- lgpformula (lgpformula-class), 28
- lgpformula-class, 28
- lgpmodel, 4, 10, 24, 28, 33, 49, 50, 53, 54
- lgpmodel (lgpmodel-class), 29
- lgpmodel-class, 29
- lgpr (lgpr-package), 3
- lgpr-package, 3
- lgprhs, 25, 26, 28, 30, 32, 34
- lgprhs (lgprhs-class), 30
- lgprhs-class, 30
- lgpscaling, 8, 11, 15, 30
- lgpscaling (lgpscaling-class), 30
- lgpscaling-class, 30
- lgpsim, 31, 45, 63
- lgpsim (lgpsim-class), 31
- lgpsim-class, 31
- lgpterm, 25, 26, 28, 30, 32, 34
- lgpterm (lgpterm-class), 32
- lgpterm-class, 32
- log_normal (priors), 48
- log_normal, (priors), 48

- mcmc_areas, 40
- mcmc_dens, 40
- mcmc_intervals, 40
- mcmc_trace, 40
- model_summary, 32

- new_x, 4-7, 33, 46, 64
- normal (priors), 48
- normal, (priors), 48
- num_components (s4_generics), 52
- num_components, GaussianPrediction-method
(GaussianPrediction-class), 18
- num_components, KernelComputer-method
(KernelComputer-class), 22
- num_components, lgpfit-method
(lgpfit-class), 27
- num_components, lgpmodel-method
(lgpmodel-class), 29
- num_components, Prediction-method
(Prediction-class), 47
- num_evalpoints (s4_generics), 52
- num_evalpoints, GaussianPrediction-method
(GaussianPrediction-class), 18
- num_evalpoints, KernelComputer-method
(KernelComputer-class), 22
- num_evalpoints, Prediction-method
(Prediction-class), 47
- num_paramsets (s4_generics), 52
- num_paramsets, GaussianPrediction-method
(GaussianPrediction-class), 18
- num_paramsets, KernelComputer-method
(KernelComputer-class), 22
- num_paramsets, Prediction-method
(Prediction-class), 47

- operations, 26, 28, 30, 32, 34
- optimize_model (sample_model), 54
- optimizing, 25, 54, 55

- param_summary (model_summary), 32
- parameter_info (s4_generics), 52
- parameter_info, lgpmodel-method
(lgpmodel-class), 29
- plot, lgpfit, missing-method
(lgpfit-class), 27
- plot, lgpsim, missing-method
(lgpsim-class), 31
- plot_api_c, 34, 37, 38, 44, 45
- plot_api_g, 35, 36, 44, 45
- plot_beta, 28
- plot_beta (plot_draws), 40
- plot_components, 4, 37, 41, 44
- plot_data, 4, 39
- plot_draws, 4, 28, 38, 40, 44
- plot_effect_times, 28
- plot_effect_times (plot_draws), 40
- plot_f, 37, 38
- plot_f (plot_pred), 43
- plot_inputwarp, 41
- plot_invgamma, 16, 42, 49
- plot_pred, 4, 38, 41, 43
- plot_sim, 31, 44
- plot_warp, 28
- plot_warp (plot_draws), 40
- postproc (s4_generics), 52
- postproc, lgpfit-method (lgpfit-class),
27
- pp_check, 45
- ppc, 45
- pred, 4, 10, 17, 20, 26, 38, 43, 46, 50, 55
- Prediction, 17, 19, 20, 38, 43, 46, 47, 50, 54
- Prediction (Prediction-class), 47
- Prediction-class, 47
- prior_pred, 4, 10, 17, 20, 26, 45, 49, 55

prior_to_num, 50
priors, 16, 25, 42, 48
qinvgamma_stanlike
 (dinvgamma_stanlike), 16
read_proteomics_data, 51, 65
relevances, 4, 52, 56
rstan, 3
s4_generics, 52
sample_model, 4, 10, 17, 20, 23, 26, 47, 50, 54
sample_param_prior, 10, 24
sample_param_prior (prior_pred), 49
sampling, 25, 49, 50, 54
select, 55
select_freq (select), 55
show, 56
show, GaussianPrediction-method
 (GaussianPrediction-class), 18
show, KernelComputer-method
 (KernelComputer-class), 22
show, lgpfite-method (lgpfite-class), 27
show, lgpfite-method (show), 56
show, lgpmodel-method (lgpmodel-class),
 29
show, lgprhs-method (show), 56
show, lgpsim-method (lgpsim-class), 31
show, lgpterm-method (show), 56
show, Prediction-method
 (Prediction-class), 47
sim.create_f, 57
sim.create_x, 57, 58
sim.create_y, 59
sim.kernels, 60
simulate_data, 61
split, 5–7, 33, 63
split_by_factor (split), 63
split_data (split), 63
split_random (split), 63
split_within_factor (split), 63
split_within_factor_random (split), 63
stanfit, 28, 50
student_t (priors), 48
student_t, (priors), 48
testdata_001, 64, 65
testdata_002, 65, 65
uniform (priors), 48
uniform, (priors), 48
validate, 66
validate_GaussianPrediction (validate),
 66
validate_lgpexpr (validate), 66
validate_lgpfit (validate), 66
validate_lgpformula (validate), 66
validate_lgpformula (validate), 66
validate_Prediction (validate), 66
var_mask, 66, 67
warp_input, 67, 67