

# Package ‘inlabru’

July 9, 2025

**Type** Package

**Title** Bayesian Latent Gaussian Modelling using INLA and Extensions

**Version** 2.13.0

**URL** <http://www.inlabru.org>, <https://inlabru-org.github.io/inlabru/>,  
<https://github.com/inlabru-org/inlabru>

**BugReports** <https://github.com/inlabru-org/inlabru/issues>

**Description** Facilitates spatial and general latent Gaussian modeling using integrated nested Laplace approximation via the INLA package (<<https://www.r-inla.org>>). Additionally, extends the GAM-like model class to more general nonlinear predictor expressions, and implements a log Gaussian Cox process likelihood for modeling univariate and spatial point processes based on ecological survey data. Model components are specified with general inputs and mapping methods to the latent variables, and the predictors are specified via general R expressions, with separate expressions for each observation likelihood model in multi-likelihood models. A prediction method based on fast Monte Carlo sampling allows posterior prediction of general expressions of the latent variables. Ecology-focused introduction in Bachl, Lindgren, Borchers, and Illian (2019) <<doi:10.1111/2041-210X.13168>>.

**License** GPL (>= 2)

**Additional\_repositories** <https://inla.r-inla-download.org/R/testing>

**RoxxygenNote** 7.3.2

**Encoding** UTF-8

**Depends** fmesher (>= 0.4.0), methods, R (>= 3.6), stats

**Imports** dplyr, lifecycle, magrittr, MatrixModels, Matrix, plyr, rlang, sf, tibble, utils, withr

**Suggests** covr, ggplot2, graphics, INLA (>= 23.01.31), knitr, maps, mgcv, patchwork, raster, RColorBrewer, rgl, rmarkdown, scales, scoringRules, shiny, sn, sp (>= 2.1), spatstat.geom, spatstat.data, sphereplot, splancs, terra, tidyterra, testthat (>= 3.2.0), tidyr, DiagrammeR

**Enhances** stars

**Config/testthat/parallel** true

**Config/testthat/edition** 3

**Collate** '0\_inlabru\_envir.R' 'bru.gof.R' 'bru.inference.R'  
 'bru\_conversion.R' 'bru\_index.R' 'bru\_input.R' 'bru\_sp.R'  
 'data.Poisson1\_1D.R' 'data.Poisson2\_1D.R' 'data.Poisson3\_1D.R'  
 'data.gorillas.R' 'data.mexdolphin.R' 'data.mrsea.R'  
 'data.robins\_subset.R' 'data.shrimp.R' 'data.toygroups.R'  
 'data.toypoints.R' 'deltaIC.R' 'deprecated.R' 'effect.R'  
 'environment.R' 'fmesher.R' 'ggplot.R' 'inla.R'  
 'inlabru-package.R' 'local\_testthat.R' 'mappers.R'  
 'mapper\_collect.R' 'mapper\_repeat.R' 'mapper\_sum.R' 'model.R'  
 'nlinla.R' 'plotsample.R' 'rgl.R' 'sampling.R' 'spatstat.R'  
 'spde.R' 'stack.R' 'track\_plotting.R' 'transformation.R'  
 'used.R' 'utils.R'

**VignetteBuilder** knitr

**BuildVignettes** true

**LazyData** true

**LazyDataCompression** xz

**NeedsCompilation** no

**Author** Finn Lindgren [aut, cre, cph] (ORCID:

<<https://orcid.org/0000-0002-5833-2011>>, Finn Lindgren continued development of the main code),

Fabian E. Bachl [aut, cph] (Fabian Bachl wrote the main code),

David L. Borchers [ctb, dtc, cph] (David Borchers wrote code for Gorilla data import and sampling, multiplot tool),

Daniel Simpson [ctb, cph] (Daniel Simpson wrote the basic LGCP sampling method),

Lindesay Scott-Howard [ctb, dtc, cph] (Lindesay Scott-Howard provided MRSea data import code),

Seaton Andy [ctb] (Andy Seaton provided testing, bugfixes, and vignettes),

Suen Man Ho [ctb, cph] (Man Ho Suen contributed features for aggregated responses and vignette updates),

Roudier Pierre [ctb, cph] (Pierre Roudier contributed general quantile summaries),

Meehan Tim [ctb, cph] (Tim Meehan contributed the SVC vignette and robins data),

Reddy Peddinenikalva Niharika [ctb, cph] (Niharika Peddinenikalva contributed the LGCP residuals vignette),

Perepolkin Dmytro [ctb, cph] (Dmytro Perepolkin contributed the ZIP/ZAP vignette)

**Maintainer** Finn Lindgren <finn.lindgren@gmail.com>

**Repository** CRAN

**Date/Publication** 2025-07-09 21:20:02 UTC

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inlabru-package      *inlabru*

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## Description

Convenient model fitting using (iterated) INLA.

## Details

`inlabru` facilitates Bayesian spatial modelling using integrated nested Laplace approximations. It is heavily based on R-inla (<https://www.r-inla.org>) but adds additional modelling abilities and simplified syntax for (in particular) spatial models. Tutorials and more information can be found at <https://inlabru-org.github.io/inlabru/> and <http://www.inlabru.org/>. The iterative method used for non-linear predictors is documented in the `method` vignette.

The main function for inference using `inlabru` is `bru()`. The general model specification details is documented in `bru_comp()` and `bru_obs()`. Posterior quantities beyond the basic summaries can be calculated with a `predict()` method, documented in `predict.bru()`. For point process inference `lgcp()` can be used as a shortcut to `bru(..., bru_obs(model="cp", ...))`.

The package comes with multiple real world data sets, namely `gorillas`, `gorillas_sf`, `mexdolphin_sf`. Plotting these data sets is straight forward using `inlabru`'s extensions to `ggplot2`, e.g. the `gg()` function. For educational purposes some simulated data sets are available as well, e.g. `Poisson1_1D`, `Poisson2_1D`, `Poisson2_1D` and `toygroups`.

## Author(s)

Fabian E. Bachl <[bachlfab@gmail.com](mailto:bachlfab@gmail.com)> and Finn Lindgren <[finn.lindgren@gmail.com](mailto:finn.lindgren@gmail.com)>

## See Also

Useful links:

- <http://www.inlabru.org>
- <https://inlabru-org.github.io/inlabru/>
- <https://github.com/inlabru-org/inlabru>
- Report bugs at <https://github.com/inlabru-org/inlabru/issues>

as_bru_mapper	<i>Methods for mapper extraction</i>
---------------	--------------------------------------

## Description

Extract a mapper from another object

## Usage

```
as_bru_mapper(x)

## S3 method for class 'bru_mapper'
as_bru_mapper(x)

## S3 method for class 'bru_comp'
as_bru_mapper(x)

## S3 method for class 'bru_subcomp'
as_bru_mapper(x)
```

## Arguments

x                   Object to convert/extract

## Value

A bru\_mapper object

## Examples

```
# Extract a mapper from a `bru_subcomp` object
as_bru_mapper(bru_comp("x", x, mapper = bm_index(4))$main)
```

bincount	<i>1D LGCP bin count simulation and comparison with data</i>
----------	--

## Description

A common procedure of analyzing the distribution of 1D points is to chose a binning and plot the data's histogram with respect to this binning. This function compares the counts that the histogram calculates to simulations from a 1D log Gaussian Cox process conditioned on the number of data samples. For each bin this results in a median number of counts as well as a confidence interval. If the LGCP is a plausible model for the observed points then most of the histogram counts (number of points within a bin) should be within the confidence intervals. Note that a proper comparison is a multiple testing problem which the function does not solve for you.

**Usage**

```
bincount(
  result,
  predictor,
  observations,
  breaks,
  nint = 20,
  probs = c(0.025, 0.5, 0.975),
  ...
)
```

**Arguments**

<code>result</code>	A result object from a <code>bru()</code> or <code>lgcp()</code> call
<code>predictor</code>	A formula describing the prediction of a 1D LGCP via <code>predict()</code> .
<code>observations</code>	A vector of observed values
<code>breaks</code>	A vector of bin boundaries
<code>nint</code>	Number of integration intervals per bin. Increase this if the bins are wide and the LGCP is not smooth.
<code>probs</code>	numeric vector of probabilities with values in [0,1]
<code>...</code>	arguments passed on to <code>predict.bru()</code>

**Value**

An `data.frame` with a `ggplot` attribute `gpp`

**Examples**

```
## Not run:
if (require(ggplot2) && require(fmesher) && bru_safe_inla()) {
  # Load a point pattern
  data(Poisson2_1D)

  # Take a look at the point (and frequency) data

  ggplot(pts2) +
    geom_histogram(
      aes(x = x),
      binwidth = 55 / 20,
      boundary = 0,
      fill = NA,
      color = "black"
    ) +
    geom_point(aes(x), y = 0, pch = "|", cex = 4) +
    coord_fixed(ratio = 1)

  # Fit an LGCP model
  x <- seq(0, 55, length.out = 50)
```

```

mesh1D <- fm_mesh_1d(x, boundary = "free")
matern <- INLA::inla.spde2.pcmatern(mesh1D,
  prior.range = c(1, 0.01),
  prior.sigma = c(1, 0.01),
  constr = TRUE
)
mdl <- x ~ spde1D(x, model = matern) + Intercept(1)
fit.spde <- lgcp(mdl, pts2, domain = list(x = mesh1D))

# Calculate bin statistics
bc <- bincount(
  result = fit.spde,
  observations = pts2,
  breaks = seq(0, max(pts2), length.out = 12),
  predictor = x ~ exp(spde1D + Intercept)
)

# Plot them!
attributes(bc)$ggp
}

## End(Not run)

```

**bm\_aggregate***Mapper for aggregation***Description**

Constructs a mapper that aggregates elements of the input state, so it can be used e.g. for weighted summation or integration over blocks of values.

**Usage**

```

bm_aggregate(rescale = FALSE, n_block = NULL, type = NULL)

bru_mapper_aggregate(...)

## S3 method for class 'bm_aggregate'
ibm_n(mapper, ..., input = NULL, state = NULL, n_state = NULL)

## S3 method for class 'bm_aggregate'
ibm_n_output(mapper, input = NULL, ...)

## S3 method for class 'bm_aggregate'
ibm_values(mapper, ..., state = NULL, n_state = NULL)

## S3 method for class 'bm_aggregate'
ibm_jacobian(mapper, input, state = NULL, ...)

```

```
## S3 method for class 'bm_aggregate'
ibm_eval(mapper, input, state = NULL, ..., sub_lin = NULL)
```

## Arguments

rescale	logical; For <code>bm_aggregate</code> and <code>bm_logsumexp</code> , specifies if the blockwise sums should be normalised by the blockwise weight sums or not:
	<ul style="list-style-type: none"> <li>• FALSE: (default) Straight weighted sum, no rescaling.</li> <li>• TRUE: Divide by the sum of the weight values within each block. This is useful for integration averages, when the given weights are plain integration weights. If the weights are <code>NULL</code> or all ones, this is the same as dividing by the number of entries in each block.</li> </ul>
n_block	Predetermined number of output blocks. If <code>NULL</code> , overrides the maximum block index in the inputs. The priority order is <code>input\$n_block</code> , the mapper definition <code>n_block</code> , then <code>max(input\$block)</code> .
type	character; if non- <code>NULL</code> , overrides the <code>rescale</code> argument, and constructs an aggregation mapper of the given type instead. Supported values are "sum", "average", "logsumexp", and "logaverageexp".
...	Arguments passed on to other methods
mapper	A mapper S3 object, inheriting from <code>bru_mapper</code> .
input	Data input for the mapper.
state	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
n_state	integer giving the length of the state vector for mappers that have state dependent output size.
sub_lin	Internal, optional pre-computed sub-mapper information

## Methods (by generic)

- `ibm_jacobian(bm_aggregate)`: `input` should be a list with elements `block` and `weights`. `block` should be a vector of the same length as the `state`, or `NULL`, with `NULL` equivalent to all-1. If `weights` is `NULL`, it's interpreted as all-1.

## See Also

`bru_mapper`, `bru_mapper_generics`

Other mappers: `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesher()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_multi()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

## Examples

```
m <- bm_aggregate()
ibm_eval2(m, list(block = c(1, 2, 1, 2), weights = 1:4), 11:14)
ibm_eval2(m, list(block = c(1, 2, 1, 2), weights = 1:4, n_block = 3), 11:14)
```

**bm\_collect***Mapper for concatenated variables***Description**

Constructs a concatenated collection mapping

**Usage**

```
bm_collect(mappers, hidden = FALSE)

bru_mapper_collect(...)

## S3 method for class 'bm_collect'
ibm_n(mapper, inla_f = FALSE, multi = FALSE, ...)

## S3 method for class 'bm_collect'
ibm_n_output(mapper, input, state = NULL, inla_f = FALSE, multi = FALSE, ...)

## S3 method for class 'bm_collect'
ibm_values(mapper, inla_f = FALSE, multi = FALSE, ...)

## S3 method for class 'bm_collect'
ibm_is_linear(mapper, inla_f = FALSE, multi = FALSE, ...)

## S3 method for class 'bm_collect'
ibm_jacobian(
  mapper,
  input,
  state = NULL,
  inla_f = FALSE,
  multi = FALSE,
  ...,
  sub_lin = NULL
)

## S3 method for class 'bm_collect'
ibm_eval(
  mapper,
  input,
  state,
  inla_f = FALSE,
  multi = FALSE,
  ...,
  sub_lin = NULL
)
```

```

## S3 method for class 'bm_collect'
ibm_linear(mapper, input, state, inla_f = FALSE, ...)

## S3 method for class 'bm_collect'
ibm_invalid_output(mapper, input, state, inla_f = FALSE, multi = FALSE, ...)

## S3 method for class 'bm_collect'
x[i, drop = TRUE]

## S3 method for class 'bru_mapper_collect'
x[i, drop = TRUE]

## S3 method for class 'bm_collect'
ibm_names(mapper)

## S3 replacement method for class 'bm_collect'
ibm_names(mapper) <- value

## S3 replacement method for class 'bru_mapper_collect'
ibm_names(mapper) <- value

```

## Arguments

<code>mappers</code>	A list of <code>bru_mapper</code> objects
<code>hidden</code>	logical, set to TRUE to flag that the mapper is to be used as a first level input mapper for <code>INLA::f()</code> in a model that requires making only the first mapper visible to <code>INLA::f()</code> and <code>INLA::inla.stack()</code> , such as for "bym2" models, as activated by the <code>inla_f</code> argument to <code>ibm_n</code> , <code>ibm_values</code> , and <code>ibm_jacobian</code> . Set to FALSE to always access the full mapper, e.g. for <code>rgeneric</code> models
<code>...</code>	Arguments passed on to other methods
<code>mapper</code>	A mapper S3 object, inheriting from <code>bru_mapper</code> .
<code>inla_f</code>	logical; when TRUE for <code>ibm_n()</code> and <code>ibm_values()</code> , the result must be compatible with the <code>INLA::f(...)</code> and corresponding <code>INLA::inla.stack(...)</code> constructions. For <code>ibm_{eval,jacobian,linear}</code> , the input interpretation may be different. Implementations do not normally need to do anything different, except for mappers of the type needed for hidden multicomponent models such as "bym2", which can be handled by <code>bm_collect</code> .
<code>multi</code>	logical; If TRUE (or positive), recurse one level into sub-mappers
<code>input</code>	Data input for the mapper.
<code>state</code>	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
<code>sub_lin</code>	Internal, optional pre-computed sub-mapper information
<code>x</code>	object from which to extract element(s)
<code>i</code>	indices specifying element(s) to extract

drop	logical; For <code>[.bm_collect</code> , whether to extract an individual mapper when <code>i</code> identifies a single element. If FALSE, a list of sub-mappers is returned (suitable e.g. for creating a new <code>bm_collect</code> object). Default: TRUE
value	a character vector of the same length as the number of sub-mappers in the mapper

### Value

- `[`-indexing a `bm_collect` extracts a subset `bm_collect` object (for `drop` FALSE) or an individual sub-mapper (for `drop` TRUE, and `i` identifies a single element)
- The `names()` method for `bm_collect` returns the names from the sub-mappers list

### Methods (by generic)

- `ibm_jacobian(bm_collect)`: Accepts a list with named entries, or a list with unnamed but ordered elements. The names must match the sub-mappers, see `ibm_names.bm_collect()`. Each list element should take a format accepted by the corresponding sub-mapper. In case each element is a vector, the input can be given as a data.frame with named columns, a matrix with named columns, or a matrix with unnamed but ordered columns. When `inla_f=TRUE` and `hidden=TRUE` in the mapper definition, the input format should instead match that of the first, non-hidden, sub-mapper.
- `ibm_invalid_output(bm_collect)`: Accepts a list with named entries, or a list with unnamed but ordered elements. The names must match the sub-mappers, see `ibm_names.bm_collect()`. Each list element should take a format accepted by the corresponding sub-mapper. In case each element is a vector, the input can be given as a data.frame with named columns, a matrix with named columns, or a matrix with unnamed but ordered columns.

### See Also

`bru_mapper`, `bru_mapper_generics`

Other mappers: `bm_aggregate()`, `bm_const()`, `bm_factor()`, `bm_fmesh()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_multi()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

### Examples

```
(m <- bm_collect(list(
  a = bm_index(2),
  b = bm_index(3)
), hidden = FALSE))
ibm_eval2(m, list(a = c(1, 2), b = c(1, 3, 2)), 1:5)
```

---

<code>bm_const</code>	<i>Constant mapper</i>
-----------------------	------------------------

---

## Description

Create a constant mapper

## Usage

```
bm_const()
bru_mapper_const()

## S3 method for class 'bm_const'
ibm_n(mapper, ...)

## S3 method for class 'bm_const'
ibm_values(mapper, ...)

## S3 method for class 'bm_const'
ibm_jacobian(mapper, input, ...)

## S3 method for class 'bm_const'
ibm_eval(mapper, input, state = NULL, ...)
```

## Arguments

mapper	A mapper S3 object, inheriting from <code>bru_mapper</code> .
...	Arguments passed on to other methods
input	Data input for the mapper.
state	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>

## See Also

`bru_mapper`, `bru_mapper_generics`

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_factor()`, `bm_fmesh()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_multi()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

## Examples

```
m <- bm_const()
ibm_eval2(m, input = 1:4)
```

---

bm_factor	<i>Mapper for factor variables</i>
-----------	------------------------------------

---

## Description

Create a factor mapper

## Usage

```
bm_factor(values, factor_mapping, indexed = FALSE)

bru_mapper_factor(...)

## S3 method for class 'bm_factor'
ibm_n(mapper, ...)

## S3 method for class 'bm_factor'
ibm_values(mapper, ...)

## S3 method for class 'bm_factor'
ibm_jacobian(mapper, input, ...)
```

## Arguments

<code>values</code>	Input values calculated by <a href="#">bru_input.bru_input()</a>
<code>factor_mapping</code>	character; selects the type of factor mapping. <ul style="list-style-type: none"> <li>• 'contrast' for leaving out the first factor level.</li> <li>• 'full' for keeping all levels.</li> </ul>
<code>indexed</code>	logical; if TRUE, the <code>ibm_values()</code> method will return an integer vector instead of the factor levels. This is needed e.g. for group and replicate mappers, since <code>INLA::f()</code> doesn't accept factor values. Default: FALSE, which works for the main input mappers. The default mapper constructions will set it the required setting.
<code>...</code>	Arguments passed on to other methods
<code>mapper</code>	A mapper S3 object, inheriting from <code>bru_mapper</code> .
<code>input</code>	Data input for the mapper.

## See Also

[bru\\_mapper](#), [bru\\_mapper\\_generics](#)

Other mappers: [bm\\_aggregate\(\)](#), [bm\\_collect\(\)](#), [bm\\_const\(\)](#), [bm\\_fmesh\(\)](#), [bm\\_harmonics\(\)](#), [bm\\_index\(\)](#), [bm\\_linear\(\)](#), [bm\\_logsumexp\(\)](#), [bm\\_marginal\(\)](#), [bm\\_matrix\(\)](#), [bm\\_mesh\\_B\(\)](#), [bm\\_multi\(\)](#), [bm\\_pipe\(\)](#), [bm\\_repeat\(\)](#), [bm\\_scale\(\)](#), [bm\\_shift\(\)](#), [bm\\_sum\(\)](#), [bm\\_taylor\(\)](#), [bru\\_get\\_mapper\(\)](#), [bru\\_mapper\(\)](#), [bru\\_mapper.fm\\_mesh\\_1d\(\)](#), [bru\\_mapper.fm\\_mesh\\_2d\(\)](#), [bru\\_mapper\\_generics](#)

## Examples

```
m <- bm_factor(factor(c("a", "b")), "full")
ibm_eval2(m, input = c("b", "a", "a", "b"), state = c(1, 3))

m <- bm_factor(factor(c("a", "b")), "contrast")
ibm_eval2(m, input = factor(c("b", "a", "a", "b")), state = 2)
```

bm\_fmesher

*Mapper for general fmesher function space objects*

## Description

Creates a mapper for general fmesher function space objects.

## Usage

```
bm_fmesher(mesh)

bru_mapper_fmesher(...)

## S3 method for class 'bm_fmesher'
ibm_n(mapper, ...)

## S3 method for class 'bm_fmesher'
ibm_values(mapper, ...)

## S3 method for class 'bm_fmesher'
ibm_jacobian(mapper, input, ...)
```

## Arguments

<code>mesh</code>	An fmesher object to map, supported by <a href="#">fmesher::fm_basis</a> ( <code>mesh, input</code> ) and <a href="#">fmesher::fm_dof</a> ( <code>mesh</code> ).
<code>...</code>	Arguments passed on to other methods
<code>mapper</code>	A mapper S3 object, inheriting from <code>bru_mapper</code> .
<code>input</code>	Data input for the mapper.

## Details

Handles indexed mapping for all fmesher classes that support `fm_dof()` and `fm_basis()` methods. For non-indexed mapping of `fm_mesh_1d` objects, use `bru_mapper(mesh, indexed = FALSE)` which invokes the [bru\\_mapper.fm\\_mesh\\_1d](#)() method.

## Value

A `bm_fmesher` object.

**See Also**

[bru\\_mapper](#), [bru\\_mapper\\_generics](#)

Other mappers: [bm\\_aggregate\(\)](#), [bm\\_collect\(\)](#), [bm\\_const\(\)](#), [bm\\_factor\(\)](#), [bm\\_harmonics\(\)](#), [bm\\_index\(\)](#), [bm\\_linear\(\)](#), [bm\\_logsumexp\(\)](#), [bm\\_marginal\(\)](#), [bm\\_matrix\(\)](#), [bm\\_mesh\\_B\(\)](#), [bm\\_multi\(\)](#), [bm\\_pipe\(\)](#), [bm\\_repeat\(\)](#), [bm\\_scale\(\)](#), [bm\\_shift\(\)](#), [bm\\_sum\(\)](#), [bm\\_taylor\(\)](#), [bru\\_get\\_mapper\(\)](#), [bru\\_mapper\(\)](#), [bru\\_mapper.fm\\_mesh\\_1d\(\)](#), [bru\\_mapper.fm\\_mesh\\_2d\(\)](#), [bru\\_mapper\\_generics](#)

**Examples**

```
m <- bm_fmeshers(fmexample$mesh)
ibm_n(m)
ibm_eval(m, as.matrix(expand.grid(-2:2, -2:2)), seq_len(ibm_n(m)))
```

**bm\_harmonics**

*Mapper for cos/sin functions*

**Description**

Constructs a mapper for cos/sin functions of orders 1 (if intercept is TRUE, otherwise 0) through order. The total number of basis functions is intercept + 2 \* order.

Optionally, each order can be given a non-unit scaling, via the scaling vector, of length intercept + order. This can be used to give an effective spectral prior. For example, let

```
scaling = 1 / (1 + (0:4)^2)
x <- seq(0, 1, length.out = 11)
bmh1 = bm_harmonics(order = 4, interval = c(0, 1))
u1 <- ibm_eval(
  bmh1,
  input = x,
  state = rnorm(9, sd = rep(scaling, c(1, 2, 2, 2))))
)
```

Then, with

```
bmh2 = bm_harmonics(order = 4, scaling = scaling)
u2 = ibm_eval(bmh2, input = x, state = rnorm(9))
```

the stochastic properties of u1 and u2 will be the same, with scaling^2 determining the variance for each frequency contribution.

The period for the first order harmonics is shifted and scaled to match interval.

## Usage

```
bm_harmonics(order = 1, scaling = 1, intercept = TRUE, interval = c(0, 1))

bru_mapper_harmonics(...)

## S3 method for class 'bm_harmonics'
ibm_n(mapper, inla_f = FALSE, ...)

## S3 method for class 'bm_harmonics'
ibm_jacobian(mapper, input, state = NULL, inla_f = FALSE, ...)
```

## Arguments

order	For <code>bm_harmonics</code> , specifies the maximum cos/sin order. (Default 1)
scaling	For <code>bm_harmonics</code> , specifies an optional vector of scaling factors of length <code>intercept + order</code> , or a common single scalar.
intercept	logical; For <code>bm_harmonics</code> , if TRUE, the first basis function is a constant. (Default TRUE)
interval	numeric length-2 vector specifying a domain interval. Default <code>c(0, 1)</code> .
...	Arguments passed on to other methods
mapper	A mapper S3 object, inheriting from <code>bru_mapper</code> .
inla_f	logical; when TRUE for <code>ibm_n()</code> and <code>ibm_values()</code> , the result must be compatible with the <code>INLA::f(...)</code> and corresponding <code>INLA::inla.stack(...)</code> constructions. For <code>ibm_{eval,jacobian,linear}</code> , the input interpretation may be different. Implementations do not normally need to do anything different, except for mappers of the type needed for hidden multicomponent models such as "bym2", which can be handled by <code>bm_collect</code> .
input	Data input for the mapper.
state	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>

## See Also

[bru\\_mapper](#), [bru\\_mapper\\_generics](#)

Other mappers: [bm\\_aggregate\(\)](#), [bm\\_collect\(\)](#), [bm\\_const\(\)](#), [bm\\_factor\(\)](#), [bm\\_fmesh\(\)](#), [bm\\_index\(\)](#), [bm\\_linear\(\)](#), [bm\\_logsumexp\(\)](#), [bm\\_marginal\(\)](#), [bm\\_matrix\(\)](#), [bm\\_mesh\\_B\(\)](#), [bm\\_multi\(\)](#), [bm\\_pipe\(\)](#), [bm\\_repeat\(\)](#), [bm\\_scale\(\)](#), [bm\\_shift\(\)](#), [bm\\_sum\(\)](#), [bm\\_taylor\(\)](#), [bru\\_get\\_mapper\(\)](#), [bru\\_mapper\(\)](#), [bru\\_mapper.fm\\_mesh\\_1d\(\)](#), [bru\\_mapper.fm\\_mesh\\_2d\(\)](#), [bru\\_mapper\\_generics](#)

## Examples

```
m <- bm_harmonics(2)
ibm_eval2(m, input = c(0, pi / 4, pi / 2, 3 * pi / 4), 1:5)
```

**bm\_index***Mapper for indexed variables*

## Description

Create a an indexing mapper

## Usage

```
bm_index(n = 1L, ...)
bru_mapper_index(...)

## S3 method for class 'bm_index'
ibm_invalid_output(mapper, input, state, ...)

## S3 method for class 'bm_index'
ibm_jacobian(mapper, input, state, ...)
```

## Arguments

<code>n</code>	Size of a model for <code>bm_index</code>
<code>...</code>	Arguments passed on to other methods
<code>mapper</code>	A mapper S3 object, inheriting from <code>bru_mapper</code> .
<code>input</code>	Data input for the mapper.
<code>state</code>	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>

## See Also

`bru_mapper`, `bru_mapper_generics`

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesh()`, `bm_harmonics()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_multi()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

## Examples

```
m <- bm_index(4)
ibm_eval(m, -2:6, 1:4)
```

---

bm_linear	<i>Mapper for a linear effect</i>
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---

## Description

Create a mapper for linear effects

## Usage

```
bm_linear()  
  
bru_mapper_linear()  
  
## S3 method for class 'bm_linear'  
ibm_n(mapper, ...)  
  
## S3 method for class 'bm_linear'  
ibm_values(mapper, ...)  
  
## S3 method for class 'bm_linear'  
ibm_jacobian(mapper, input, ...)
```

## Arguments

mapper	A mapper S3 object, inheriting from bru_mapper.
...	Arguments passed on to other methods
input	Data input for the mapper.

## See Also

[bru\\_mapper](#), [bru\\_mapper\\_generics](#)

Other mappers: [bm\\_aggregate\(\)](#), [bm\\_collect\(\)](#), [bm\\_const\(\)](#), [bm\\_factor\(\)](#), [bm\\_fmesh\(\)](#), [bm\\_harmonics\(\)](#), [bm\\_index\(\)](#), [bm\\_logsumexp\(\)](#), [bm\\_marginal\(\)](#), [bm\\_matrix\(\)](#), [bm\\_mesh\\_B\(\)](#), [bm\\_multi\(\)](#), [bm\\_pipe\(\)](#), [bm\\_repeat\(\)](#), [bm\\_scale\(\)](#), [bm\\_shift\(\)](#), [bm\\_sum\(\)](#), [bm\\_taylor\(\)](#), [bru\\_get\\_mapper\(\)](#), [bru\\_mapper\(\)](#), [bru\\_mapper.fm\\_mesh\\_1d\(\)](#), [bru\\_mapper.fm\\_mesh\\_2d\(\)](#), [bru\\_mapper\\_generics](#)

## Examples

```
m <- bm_linear()  
ibm_eval(m, input = 1:4, state = 2)
```

---

 bm\_list *Methods for mapper lists*


---

### Description

`bru_mapper` lists can be combined into `bm_list` lists.

### Usage

```
as_bm_list(x)

## S3 method for class 'list'
as_bm_list(x)

## S3 method for class 'bm_list'
as_bm_list(x)

## S3 method for class 'bru_comp_list'
as_bm_list(x)

## S3 method for class 'bru_mapper'
c(...)

## S3 method for class 'bm_list'
c(...)

## S3 method for class 'bm_list'
x[i]

## S3 method for class 'bm_list'
ibm_linear(mapper, input, state = NULL, ...)

## S3 method for class 'bm_list'
ibm_simplify(mapper, input = NULL, state = NULL, ...)
```

### Arguments

<code>x</code>	<code>bm_list</code> object from which to extract element(s)
<code>...</code>	Objects to be combined.
<code>i</code>	indices specifying elements to extract
<code>mapper</code>	A mapper S3 object, inheriting from <code>bru_mapper</code> .
<code>input</code>	Data input for the mapper.
<code>state</code>	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>

**Value**

A `bm_list` object

**Methods (by generic)**

- `c(bm_list)`: The ... arguments should be `bm_list` objects.
- `[:`: Extract sub-list

**Functions**

- `c(bru_mapper)`: The ... arguments should be `bru_mapper` objects.

**Examples**

```
m <- c(A = bm_const(), B = bm_scale())
str(m)
str(m[2])
```

`bm_logsumexp`

*Mapper for log-sum-exp aggregation*

**Description**

Constructs a mapper that aggregates elements of `exp(state)`, with optional non-negative weighting, and then takes the `log()`, so it can be used e.g. for  $v_k = \log[\sum_{i \in I_k} w_i \exp(u_i)]$  and  $v_k = \log[\sum_{i \in I_k} w_i \exp(u_i) / \sum_{i \in I_k} w_i]$  calculations. Relies on the input handling methods for `bm_aggregate`, but also allows the weights to be supplied on a logarithmic scale as `log_weights`. To avoid numerical overflow, it uses the common method of internally shifting the state blockwise;  $v_k = s_k + \log[\sum_{i \in I_k} \exp(u_i + \log(w_i) - s_k)]$ , where  $s_k = \max_{i \in I_k} u_i + \log(w_i)$  is the shift for block  $k$ .

**Usage**

```
bm_logsumexp(rescale = FALSE, n_block = NULL)

bru_mapper_logsumexp(...)

## S3 method for class 'bm_logsumexp'
ibm_jacobian(mapper, input, state = NULL, ...)

## S3 method for class 'bm_logsumexp'
ibm_eval(mapper, input, state = NULL, log = TRUE, ..., sub_lin = NULL)
```

## Arguments

<code>rescale</code>	logical; For <code>bm_aggregate</code> and <code>bm_logsumexp</code> , specifies if the blockwise sums should be normalised by the blockwise weight sums or not:
	<ul style="list-style-type: none"> <li>• FALSE: (default) Straight weighted sum, no rescaling.</li> <li>• TRUE: Divide by the sum of the weight values within each block. This is useful for integration averages, when the given weights are plain integration weights. If the weights are NULL or all ones, this is the same as dividing by the number of entries in each block.</li> </ul>
<code>n_block</code>	Predetermined number of output blocks. If NULL, overrides the maximum block index in the inputs. The priority order is <code>input\$n_block</code> , the mapper definition <code>n_block</code> , then <code>max(input\$block)</code> .
<code>...</code>	Arguments passed on to other methods
<code>mapper</code>	A mapper S3 object, inheriting from <code>bru_mapper</code> .
<code>input</code>	Data input for the mapper.
<code>state</code>	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
<code>log</code>	logical; control log output. Default TRUE, see the <code>ibm_eval()</code> details for <code>logsumexp</code> mappers.
<code>sub_lin</code>	Internal, optional pre-computed sub-mapper information

## Methods (by generic)

- `ibm_jacobian(bm_logsumexp)`: `input` should be a list with elements `block` and `weights`. `block` should be a vector of the same length as the `state`, or NULL, with NULL equivalent to all-1. If `weights` is NULL, it's interpreted as all-1.
- `ibm_eval(bm_logsumexp)`: When `log` is TRUE (default), `ibm_eval()` for `logsumexp` returns the log-sum-weight-exp value. If FALSE, the sum-weight-exp value is returned.

## See Also

[bru\\_mapper](#), [bru\\_mapper\\_generics](#)

Other mappers: [bm\\_aggregate\(\)](#), [bm\\_collect\(\)](#), [bm\\_const\(\)](#), [bm\\_factor\(\)](#), [bm\\_fmesh\(\)](#), [bm\\_harmonics\(\)](#), [bm\\_index\(\)](#), [bm\\_linear\(\)](#), [bm\\_marginal\(\)](#), [bm\\_matrix\(\)](#), [bm\\_mesh\\_B\(\)](#), [bm\\_multi\(\)](#), [bm\\_pipe\(\)](#), [bm\\_repeat\(\)](#), [bm\\_scale\(\)](#), [bm\\_shift\(\)](#), [bm\\_sum\(\)](#), [bm\\_taylor\(\)](#), [bru\\_get\\_mapper\(\)](#), [bru\\_mapper\(\)](#), [bru\\_mapper.fm\\_mesh\\_1d\(\)](#), [bru\\_mapper.fm\\_mesh\\_2d\(\)](#), [bru\\_mapper\\_generics](#)

## Examples

```
m <- bm_logsumexp()
ibm_eval2(m, list(block = c(1, 2, 1, 2), weights = 1:4), 11:14)
ibm_eval2(m, list(block = c(1, 2, 1, 2), weights = 1:4, n_block = 3), 11:14)
```

---

bm_marginal	<i>Mapper for marginal distribution transformation</i>
-------------	--

---

## Description

Constructs a mapper that transforms the marginal distribution state from  $N(0, 1)$  to the distribution of a given (continuous) quantile function. The ... arguments are used as parameter arguments to qfun, pfun, dfun, and dqfun.

## Usage

```
bm_marginal(qfun, pfun = NULL, dfun = NULL, dqfun = NULL, ..., inverse = FALSE)

bru_mapper_marginal(...)

## S3 method for class 'bm_marginal'
ibm_n(mapper, ..., state = NULL, n_state = NULL)

## S3 method for class 'bm_marginal'
ibm_n_output(mapper, input, state = NULL, ..., n_state = NULL)

## S3 method for class 'bm_marginal'
ibm_values(mapper, ..., state = NULL, n_state = NULL)

## S3 method for class 'bm_marginal'
ibm_jacobian(mapper, input, state = NULL, ..., reverse = FALSE)

## S3 method for class 'bm_marginal'
ibm_eval(mapper, input, state = NULL, ..., reverse = FALSE)
```

## Arguments

qfun	A quantile function, supporting lower.tail and log.p arguments, like <code>stats::qnorm()</code> .
pfun	A CDF, supporting lower.tail and log.p arguments, like <code>stats::pnorm()</code> . Only needed and used when xor(mapper[["inverse"]], reverse) is TRUE in a method call. Default NULL
dfun	A pdf, supporting log argument, like <code>stats::dnorm()</code> . If NULL (default), uses finite differences on qfun or pfun instead.
dqfun	A function evaluating the reciprocal of the derivative of qfun. If NULL (default), uses dfun(qfun(...), ...) or finite differences on qfun or pfun instead.
...	Arguments passed on to other methods
inverse	logical; If FALSE (default), <code>bm_marginal()</code> defines a mapping from standard Normal to a specified distribution. If TRUE, it defines a mapping from the specified distribution to a standard Normal.
mapper	A mapper S3 object, inheriting from <code>bru_mapper</code> .

state	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
n_state	integer giving the length of the state vector for mappers that have state dependent output size.
input	Data input for the mapper.
reverse	logical; control <code>bm_marginal</code> evaluation. Default FALSE. When TRUE, reverses the direction of the mapping, see details for marginal mappers.

### Methods (by generic)

- `ibm_jacobian(bm_marginal)`: Non-NULL input values are interpreted as a parameter list for `qfun`, overriding that of the mapper itself.
- `ibm_eval(bm_marginal)`: When `xor(mapper[["inverse"]], reverse)` is FALSE, `ibm_eval()` for marginal returns `qfun(pnorm(x), param)`, evaluated in a numerically stable way. Otherwise, evaluates the inverse `qnorm(pfun(x, param))` instead.

### See Also

[bru\\_mapper](#), [bru\\_mapper\\_generics](#)

Other mappers: [bm\\_aggregate\(\)](#), [bm\\_collect\(\)](#), [bm\\_const\(\)](#), [bm\\_factor\(\)](#), [bm\\_fmesh\(\)](#), [bm\\_harmonics\(\)](#), [bm\\_index\(\)](#), [bm\\_linear\(\)](#), [bm\\_logsumexp\(\)](#), [bm\\_matrix\(\)](#), [bm\\_mesh\\_B\(\)](#), [bm\\_multi\(\)](#), [bm\\_pipe\(\)](#), [bm\\_repeat\(\)](#), [bm\\_scale\(\)](#), [bm\\_shift\(\)](#), [bm\\_sum\(\)](#), [bm\\_taylor\(\)](#), [bru\\_get\\_mapper\(\)](#), [bru\\_mapper\(\)](#), [bru\\_mapper.fm\\_mesh\\_1d\(\)](#), [bru\\_mapper.fm\\_mesh\\_2d\(\)](#), [bru\\_mapper\\_generics](#)

### Examples

```
m <- bm_marginal(qexp, pexp, rate = 1 / 8)
(val <- ibm_eval(m, state = -5:5))
ibm_eval(m, state = val, reverse = TRUE)
m <- bm_marginal(qexp, pexp, dexp, rate = 1 / 8)
ibm_eval2(m, state = -3:3)
```

**bm\_matrix**

*Mapper for matrix multiplication*

### Description

Create a matrix mapper, for a given number of columns

### Usage

```
bm_matrix(labels)

bru_mapper_matrix(...)

## S3 method for class 'bm_matrix'
```

```

ibm_n(mapper, ...)

## S3 method for class 'bm_matrix'
ibm_values(mapper, ...)

## S3 method for class 'bm_matrix'
ibm_jacobian(mapper, input, state = NULL, inla_f = FALSE, ...)

```

## Arguments

labels	Column labels for matrix mappings; Can be factor, character, or a single integer specifying the number of columns for integer column indexing.
...	Arguments passed on to other methods
mapper	A mapper S3 object, inheriting from <code>bru_mapper</code> .
input	Data input for the mapper.
state	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
inla_f	logical; when TRUE for <code>ibm_n()</code> and <code>ibm_values()</code> , the result must be compatible with the <code>INLA:::f(...)</code> and corresponding <code>INLA:::inla.stack(...)</code> constructions. For <code>ibm_{eval,jacobian,linear}</code> , the input interpretation may be different. Implementations do not normally need to do anything different, except for mappers of the type needed for hidden multicomponent models such as "bym2", which can be handled by <code>bm_collect</code> .

## See Also

`bru_mapper`, `bru_mapper_generics`

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesh()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_mesh_B()`, `bm_multi()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

## Examples

```

m <- bm_matrix(labels = c("a", "b"))
ibm_values(m)
ibm_eval2(m, input = matrix(1:6, 3, 2), state = 2:3)

m <- bm_matrix(labels = 2L)
ibm_values(m)
ibm_eval2(m, input = matrix(1:6, 3, 2), state = 2:3)

```

---

<b>bm_mesh_B</b>	<i>Mapper for basis conversion</i>
------------------	------------------------------------

---

## Description

Creates a mapper for handling basis conversions

## Usage

```
bm_mesh_B(mesh, B)

bru_mapper_mesh_B(...)

## S3 method for class 'bm_mesh_B'
ibm_n(mapper, ...)

## S3 method for class 'bm_mesh_B'
ibm_values(mapper, ...)

## S3 method for class 'bm_mesh_B'
ibm_jacobian(mapper, input, ...)
```

## Arguments

mesh	object supported by <code>bru_mapper</code> , typically <code>fm_mesh_2d</code> or <code>fm_mesh_1d</code>
B	a square or tall basis conversion matrix
...	Arguments passed on to other methods
mapper	A mapper S3 object, inheriting from <code>bru_mapper</code> .
input	The values for which to produce a mapping matrix

## See Also

`bru_mapper`, `bru_mapper_generics`

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesh()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_multi()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

---

**bm\_multi***Mapper for tensor product domains*

---

## Description

Constructs a row-wise Kronecker product mapping of linear/affine mappers. Any offset in sub-mappers is added into a combined offset. Only linear/affine sub-mappers are allowed.

## Usage

```
bm_multi(mappers)

bru_mapper_multi(mappers)

## S3 method for class 'bm_multi'
ibm_n(mapper, inla_f = FALSE, multi = FALSE, ...)

## S3 method for class 'bm_multi'
ibm_n_output(mapper, input, ...)

## S3 method for class 'bm_multi'
ibm_values(mapper, inla_f = FALSE, multi = FALSE, ...)

## S3 method for class 'bm_multi'
ibm_is_linear(mapper, multi = FALSE, ...)

## S3 method for class 'bm_multi'
ibm_jacobian(
  mapper,
  input,
  state = NULL,
  inla_f = FALSE,
  multi = FALSE,
  ...,
  sub_A = NULL
)

## S3 method for class 'bm_multi'
ibm_linear(mapper, input, state, inla_f = FALSE, ...)

## S3 method for class 'bm_multi'
ibm_eval(
  mapper,
  input,
  state = NULL,
  inla_f = FALSE,
  ...,
```

```

jacobian = NULL,
pre_A = deprecated()
)

## S3 method for class 'bm_multi'
ibm_invalid_output(mapper, input, state, inla_f = FALSE, multi = FALSE, ...)

## S3 method for class 'bm_multi'
x[i, drop = TRUE]

## S3 method for class 'bru_mapper_multi'
x[i, drop = TRUE]

## S3 method for class 'bm_multi'
ibm_names(mapper)

## S3 replacement method for class 'bm_multi'
ibm_names(mapper) <- value

## S3 replacement method for class 'bru_mapper_multi'
ibm_names(mapper) <- value

```

## Arguments

<code>mappers</code>	A list of <code>bru_mapper</code> objects
<code>mapper</code>	A mapper S3 object, inheriting from <code>bru_mapper</code> .
<code>inla_f</code>	logical; when TRUE for <code>ibm_n()</code> and <code>ibm_values()</code> , the result must be compatible with the <code>INLA::f(...)</code> and corresponding <code>INLA::inla.stack(...)</code> constructions. For <code>ibm_{eval,jacobian,linear}</code> , the input interpretation may be different. Implementations do not normally need to do anything different, except for mappers of the type needed for hidden multicomponent models such as "bym2", which can be handled by <code>bm_collect</code> .
<code>multi</code>	logical; If TRUE (or positive), recurse one level into sub-mappers
<code>...</code>	Arguments passed on to other methods
<code>input</code>	Data input for the mapper.
<code>state</code>	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
<code>sub_A</code>	Internal; precomputed Jacobian matrices.
<code>jacobian</code>	For <code>ibm_eval()</code> methods, an optional pre-computed Jacobian, typically supplied by internal methods that already have the Jacobian.
<code>pre_A</code>	<b>[Deprecated]</b> in favour of <code>jacobian</code> .
<code>x</code>	object from which to extract element(s)
<code>i</code>	indices specifying element(s) to extract
<code>drop</code>	logical; For <code>[.bm_multi</code> , whether to extract an individual mapper when <code>i</code> identifies a single element. If FALSE, a list of sub-mappers is returned (suitable e.g. for creating a new <code>bm_multi</code> object). Default: TRUE

value	a character vector of up to the same length as the number of mappers in the multi-mapper x
-------	--

**Value**

- [-indexing a `bm_multi` extracts a subset `bm_multi` object (for drop FALSE) or an individual sub-mapper (for drop TRUE, and i identifies a single element)

**Methods (by generic)**

- `ibm_jacobian(bm_multi)`: Accepts a list with named entries, or a list with unnamed but ordered elements. The names must match the sub-mappers, see `ibm_names.bm_multi()`. Each list element should take a format accepted by the corresponding sub-mapper. In case each element is a vector, the input can be given as a data.frame with named columns, a matrix with named columns, or a matrix with unnamed but ordered columns.
- `ibm_invalid_output(bm_multi)`: Accepts a list with named entries, or a list with unnamed but ordered elements. The names must match the sub-mappers, see `ibm_names.bm_multi()`. Each list element should take a format accepted by the corresponding sub-mapper. In case each element is a vector, the input can be given as a data.frame with named columns, a matrix with named columns, or a matrix with unnamed but ordered columns.
- `ibm_names(bm_multi)`: Returns the names from the sub-mappers list

**See Also**

`bru_mapper`, `bru_mapper_generics`

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesh()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

**Examples**

```
(m <- bm_multi(list(a = bm_index(2), b = bm_index(3))))
ibm_eval2(m, list(a = c(1, 2, 1), b = c(1, 3, 2)), 1:6)
```

`bm_pipe`

*Mapper for linking several mappers in sequence*

**Description**

Create a pipe mapper, where `mappers` is a list of mappers, and the evaluated output of each mapper is handed as the state to the next mapper. The input format for the `ibm_eval` and `ibm_jacobian` methods is a list of inputs, one for each mapper.

## Usage

```

bm_pipe(mappers)

bru_mapper_pipe(...)

## S3 method for class 'bm_pipe'
ibm_n(mapper, ..., input = NULL, state = NULL)

## S3 method for class 'bm_pipe'
ibm_n_output(mapper, input, state = NULL, ..., n_state = NULL)

## S3 method for class 'bm_pipe'
ibm_values(mapper, ...)

## S3 method for class 'bm_pipe'
ibm_jacobian(mapper, input, state = NULL, ...)

## S3 method for class 'bm_pipe'
ibm_eval(mapper, input, state = NULL, ...)

## S3 method for class 'bm_pipe'
ibm_eval2(mapper, input, state = NULL, ...)

## S3 method for class 'bm_pipe'
ibm_simplify(
  mapper,
  input = NULL,
  state = NULL,
  inla_f = FALSE,
  ...,
  n_state = NULL
)

```

## Arguments

<code>mappers</code>	A list of <code>bru_mapper</code> objects
<code>...</code>	Arguments passed on to other methods
<code>mapper</code>	A mapper S3 object, inheriting from <code>bru_mapper</code> .
<code>input</code>	Data input for the mapper.
<code>state</code>	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
<code>n_state</code>	integer giving the length of the state vector for mappers that have state dependent output size.
<code>inla_f</code>	logical; when TRUE for <code>ibm_n()</code> and <code>ibm_values()</code> , the result must be compatible with the <code>INLA::f(...)</code> and corresponding <code>INLA::inla.stack(...)</code> constructions. For <code>ibm_{eval,jacobian,linear}</code> , the input interpretation may

be different. Implementations do not normally need to do anything different, except for mappers of the type needed for hidden multicomponent models such as "bym2", which can be handled by `bm_collect`.

### Methods (by generic)

- `ibm_simplify(bm_pipe)`: Constructs a simplified pipe mapper. For fully linear pipes, calls `ibm_linear()`. For partially non-linear pipes, replaces each sequence of linear mappers with a single `bm_taylor()` mapper, while keeping the full list of original mapper names, allowing the original input structure to be used also with the simplified mappers, since the taylor mappers are not dependent on inputs.

### See Also

`bru_mapper`, `bru_mapper_generics`

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesh()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_multi()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

### Examples

```
m <- bm_pipe(list(
  scale = bm_scale(),
  shift = bm_shift()
))
ibm_eval2(m, input = list(scale = 2, shift = 1:4), state = 1:4)
```

`bm_repeat`

*Mapper for repeating a mapper*

### Description

Defines a repeated-space mapper that sums the contributions for each copy. The `ibm_n()` method returns `ibm_n(mapper) * n_rep`, and `ibm_values()` returns `seq_len(ibm_n(mapper))`.

### Usage

```
bm_repeat(mapper, n_rep, interleaved = FALSE)

bru_mapper_repeat(...)

## S3 method for class 'bm_repeat'
ibm_n(mapper, ...)

## S3 method for class 'bm_repeat'
ibm_n_output(mapper, ...)
```

```

## S3 method for class 'bm_repeat'
ibm_values(mapper, ...)

## S3 method for class 'bm_repeat'
ibm_jacobian(
  mapper,
  input,
  state = NULL,
  inla_f = FALSE,
  multi = FALSE,
  ...,
  sub_lin = NULL
)

## S3 method for class 'bm_repeat'
ibm_eval(mapper, input, state, multi = FALSE, ..., sub_lin = NULL)

## S3 method for class 'bm_repeat'
ibm_linear(mapper, input, state, ...)

## S3 method for class 'bm_repeat'
ibm_invalid_output(mapper, input, state, ...)

```

## Arguments

<code>mapper</code>	The mapper to be repeated.
<code>n_rep</code>	The number of times to repeat the mapper. If a vector, the non-interleaved repeats are combined into a single repeat mapping, and combined with interleaved repeats via a <code>bm_sum()</code> of mappers.
<code>interleaved</code>	logical; if TRUE, the repeated mapping columns are interleaved; ( $x1[1]$ , $x2[1]$ , ..., $x1[2]$ , $x2[2]$ , ...). If FALSE (default), the repeated mapping columns are contiguous, ( $x1[1]$ , $x1[2]$ , ..., $x2[1]$ , $x2[2]$ , ...), and the Jacobian is a <code>cbind()</code> of the Jacobians of the repeated mappers. If <code>n_rep</code> is a vector, <code>interleaved</code> should either be a single logical, or a vector of the same length. Each element applies to the corresponding <code>n_rep</code> repetition specification.
<code>...</code>	Arguments passed on to other methods
<code>input</code>	Data input for the mapper.
<code>state</code>	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
<code>inla_f</code>	logical; when TRUE for <code>ibm_n()</code> and <code>ibm_values()</code> , the result must be compatible with the <code>INLA::f(...)</code> and corresponding <code>INLA::inla.stack(...)</code> constructions. For <code>ibm_{eval,jacobian,linear}</code> , the <code>input</code> interpretation may be different. Implementations do not normally need to do anything different, except for mappers of the type needed for hidden multicomponent models such as "bym2", which can be handled by <code>bm_collect</code> .
<code>multi</code>	logical; If TRUE (or positive), recurse one level into sub-mappers

sub_lin	Internal, optional pre-computed sub-mapper information
---------	--

### Value

A `bm_repeat` or `bm_sum` object, or the original input mapper.

### Methods (by generic)

- `ibm_jacobian(bm_repeat)`: The input should take the format of the repeated submapper.
- `ibm_invalid_output(bm_repeat)`: Passes on the input to the corresponding method.

### See Also

[bru\\_mapper](#), [bru\\_mapper\\_generics](#)

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesh()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_multi()`, `bm_pipe()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

### Examples

```
(m0 <- bm_index(3))
(m <- bm_repeat(m0, 5))
ibm_n(m)
ibm_values(m)
ibm_jacobian(m, 1:3)
ibm_eval(m, 1:3, seq_len(ibm_n(m)))

# Interleaving and grouping
(m <- bm_repeat(m0, c(2, 1, 2), c(TRUE, FALSE, FALSE)))
ibm_n(m)
ibm_values(m)
ibm_jacobian(m, 1:3)
ibm_eval(m, 1:3, seq_len(ibm_n(m)))
```

### Description

Create a standalone scaling mapper that can be used as part of a `bm_pipe`. If `mapper` is non-null, the `bm_scale()` constructor returns `bm_pipe(list(mapper = mapper, scale = bm_scale()))`

## Usage

```
bm_scale(mapper = NULL)

bru_mapper_scale(...)

## S3 method for class 'bm_scale'
ibm_n(mapper, ..., state = NULL, n_state = NULL)

## S3 method for class 'bm_scale'
ibm_n_output(mapper, input, state = NULL, ..., n_state = NULL)

## S3 method for class 'bm_scale'
ibm_values(mapper, ..., state = NULL, n_state = NULL)

## S3 method for class 'bm_scale'
ibm_jacobian(mapper, input, state = NULL, ..., sub_lin = NULL)

## S3 method for class 'bm_scale'
ibm_eval(mapper, input, state = NULL, ..., sub_lin = NULL)
```

## Arguments

<code>mapper</code>	For <code>bm_scale()</code> , an optional <code>bru_mapper</code> to be scaled. For <code>ibm_*</code> methods, a <code>bm_scale</code> mapper object.
<code>...</code>	Arguments passed on to other methods
<code>state</code>	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
<code>n_state</code>	integer giving the length of the state vector for mappers that have state dependent output size.
<code>input</code>	Data input for the mapper.
<code>sub_lin</code>	Internal, optional pre-computed sub-mapper information

## Methods (by generic)

- `ibm_jacobian(bm_scale)`: input `NULL` values are interpreted as no scaling.

## See Also

[bru\\_mapper](#), [bru\\_mapper\\_generics](#)

Other mappers: [bm\\_aggregate\(\)](#), [bm\\_collect\(\)](#), [bm\\_const\(\)](#), [bm\\_factor\(\)](#), [bm\\_fmesh\(\)](#), [bm\\_harmonics\(\)](#), [bm\\_index\(\)](#), [bm\\_linear\(\)](#), [bm\\_logsumexp\(\)](#), [bm\\_marginal\(\)](#), [bm\\_matrix\(\)](#), [bm\\_mesh\\_B\(\)](#), [bm\\_multi\(\)](#), [bm\\_pipe\(\)](#), [bm\\_repeat\(\)](#), [bm\\_shift\(\)](#), [bm\\_sum\(\)](#), [bm\\_taylor\(\)](#), [bru\\_get\\_mapper\(\)](#), [bru\\_mapper\(\)](#), [bru\\_mapper.fm\\_mesh\\_1d\(\)](#), [bru\\_mapper.fm\\_mesh\\_2d\(\)](#), [bru\\_mapper\\_generics](#)

## Examples

```
m <- bm_scale()
ibm_eval2(m, c(1, 2, 1, 2), 1:4)
```

**bm\_shift**

*Mapper for element-wise shifting*

## Description

Create a standalone shift mapper that can be used as part of a `bm_pipe`. If `mapper` is non-null, the `bm_shift()` constructor returns `bm_pipe(list(mapper = mapper, shift = bm_shift()))`

## Usage

```
bm_shift(mapper = NULL)

bru_mapper_shift(...)

## S3 method for class 'bm_shift'
ibm_n(mapper, ..., state = NULL, n_state = NULL)

## S3 method for class 'bm_shift'
ibm_n_output(mapper, input, state = NULL, ..., n_state = NULL)

## S3 method for class 'bm_shift'
ibm_values(mapper, ..., state = NULL, n_state = NULL)

## S3 method for class 'bm_shift'
ibm_jacobian(mapper, input, state = NULL, ..., sub_lin = NULL)

## S3 method for class 'bm_shift'
ibm_eval(mapper, input, state = NULL, ..., sub_lin = NULL)
```

## Arguments

<code>mapper</code>	A mapper S3 object, inheriting from <code>bru_mapper</code> .
<code>...</code>	Arguments passed on to other methods
<code>state</code>	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
<code>n_state</code>	integer giving the length of the state vector for mappers that have state dependent output size.
<code>input</code>	Data input for the mapper.
<code>sub_lin</code>	Internal, optional pre-computed sub-mapper information

### Methods (by generic)

- `ibm_jacobian(bm_shift)`: input NULL values are interpreted as no shift.

### See Also

`bru_mapper`, `bru_mapper_generics`

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesh()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_multi()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_sum()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

### Examples

```
m <- bm_shift()
ibm_eval2(m, c(1, 2, 1, 2), 1:4)
```

`bm_sum`

*Mapper for adding multiple mappers*

### Description

Defines a mapper that adds the effects of each submapper. The `ibm_n()` method returns the sum of `ibm_n(mappers[[k]])`, and `ibm_values()` returns `seq_len(ibm_n(mapper))`.

### Usage

```
bm_sum(mappers, single_input = FALSE)

bru_mapper_sum(...)

## S3 method for class 'bm_sum'
ibm_n(mapper, inla_f = FALSE, multi = FALSE, ...)

## S3 method for class 'bm_sum'
ibm_n_output(mapper, input, state = NULL, ...)

## S3 method for class 'bm_sum'
ibm_values(mapper, inla_f = FALSE, multi = FALSE, ...)

## S3 method for class 'bm_sum'
ibm_is_linear(mapper, multi = FALSE, ...)

## S3 method for class 'bm_sum'
ibm_jacobian(
  mapper,
  input,
```

```

state = NULL,
inla_f = FALSE,
multi = FALSE,
...,
sub_lin = NULL
)

## S3 method for class 'bm_sum'
ibm_eval(mapper, input, state, multi = FALSE, ..., sub_lin = NULL)

## S3 method for class 'bm_sum'
ibm_linear(mapper, input, state, ...)

## S3 method for class 'bm_sum'
ibm_invalid_output(mapper, input, state, multi = FALSE, ...)

## S3 method for class 'bm_sum'
x[i, drop = TRUE]

## S3 method for class 'bru_mapper_sum'
x[i, drop = TRUE]

## S3 method for class 'bm_sum'
ibm_names(mapper)

## S3 replacement method for class 'bm_sum'
ibm_names(mapper) <- value

## S3 replacement method for class 'bru_mapper_sum'
ibm_names(mapper) <- value

```

## Arguments

mappers	A list of bru_mapper objects.
single_input	logical. If TRUE, the input is passed to all sub-mappers. Otherwise, the input should be a list, data.frame, or matrix. If the mappers list has named entries, the input can reference their corresponding sub-mapper using its name.
...	Arguments passed on to other methods
mapper	A mapper S3 object, inheriting from bru_mapper.
inla_f	logical; when TRUE for ibm_n() and ibm_values(), the result must be compatible with the INLA::f(...) and corresponding INLA::inla.stack(...) constructions. For ibm_{eval,jacobian,linear}, the input interpretation may be different. Implementations do not normally need to do anything different, except for mappers of the type needed for hidden multicomponent models such as "bym2", which can be handled by bm_collect.
multi	logical; If TRUE (or positive), recurse one level into sub-mappers
input	Data input for the mapper.

state	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
sub_lin	Internal, optional pre-computed sub-mapper information
x	object from which to extract element(s)
i	indices specifying element(s) to extract
drop	logical; For <code>[ .bm_sum</code> , whether to extract an individual mapper when i identifies a single element. If FALSE, a list of sub-mappers is returned (suitable e.g. for creating a new <code>bm_sum</code> object). Default: TRUE
value	a character vector of the same length as the number of sub-mappers in the mapper

### Value

A `bm_sum` object.

- [-indexing a `bm_sum` extracts a subset `bm_sum` object (for drop FALSE) or an individual sub-mapper (for drop TRUE, and i identifies a single element)
- The `names()` method for `bm_sum` returns the names from the sub-mappers list

### Methods (by generic)

- `ibm_jacobian(bm_sum)`: Accepts a list with named entries, or a list with unnamed but ordered elements. The names must match the sub-mappers, see `ibm_names.bm_sum()`. Each list element should take a format accepted by the corresponding sub-mapper. In case each element is a vector, the input can be given as a data.frame with named columns, a matrix with named columns, or a matrix with unnamed but ordered columns.
- `ibm_invalid_output(bm_sum)`: Accepts a list with named entries, or a list with unnamed but ordered elements. The names must match the sub-mappers, see `ibm_names.bm_sum()`. Each list element should take a format accepted by the corresponding sub-mapper. In case each element is a vector, the input can be given as a data.frame with named columns, a matrix with named columns, or a matrix with unnamed but ordered columns.

### See Also

`bru_mapper`, `bru_mapper_generics`

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesh()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_multi()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

### Examples

```
(m <- bm_sum(list(a = bm_index(3), b = bm_index(2))))
ibm_n(m)
ibm_values(m)
ibm_jacobian(m, list(a = 1:3, b = c(1, 1, 2)))
ibm_eval()
```

```

  m,
  list(a = 1:3, b = c(1, 1, 2)),
  seq_len(ibm_n(m))
)

```

---

bm\_taylor*Mapper for linear Taylor approximations*

---

## Description

Provides a pre-computed affine mapping, internally used to represent and evaluate linearisation information. The `state0` information indicates for which state the offset was evaluated; The affine mapper output is defined as `effect(state) = offset + jacobian %*% (state - state0)`

## Usage

```

bm_taylor(offset = NULL, jacobian = NULL, state0 = NULL, values_mapper = NULL)

bru_mapper_taylor(...)

## S3 method for class 'bm_taylor'
ibm_n(mapper, inla_f = FALSE, multi = FALSE, ...)

## S3 method for class 'bm_taylor'
ibm_n_output(mapper, input, ...)

## S3 method for class 'bm_taylor'
ibm_values(mapper, inla_f = FALSE, multi = FALSE, ...)

## S3 method for class 'bm_taylor'
ibm_jacobian(mapper, ..., multi = FALSE)

## S3 method for class 'bm_taylor'
ibm_eval(mapper, input = NULL, state = NULL, ...)

```

## Arguments

<code>offset</code>	For <code>bm_taylor</code> , an offset vector giving the value of the linearisation at <code>state0</code> . May be <code>NULL</code> , interpreted as an all-zero vector of length determined by a non-null Jacobian.
<code>jacobian</code>	For <code>bm_taylor()</code> , the Jacobian matrix, evaluated at <code>state0</code> , or, a named list of such matrices. May be <code>NULL</code> or an empty list, for a constant mapping.
<code>state0</code>	For <code>bm_taylor</code> , the reference state for the linearisation, or a list of such states matching the <code>jacobian</code> list. <code>NULL</code> is interpreted as 0.
<code>values_mapper</code>	mapper object to be used for <code>ibm_n</code> and <code>ibm_values</code> for <code>inla_f=TRUE</code> (experimental, currently unused)

...	Arguments passed on to other methods
mapper	A mapper S3 object, inheriting from <code>bru_mapper</code> .
inla_f	logical; when TRUE for <code>ibm_n()</code> and <code>ibm_values()</code> , the result must be compatible with the <code>INLA::f(...)</code> and corresponding <code>INLA::inla.stack(...)</code> constructions. For <code>ibm_{eval,jacobian,linear}</code> , the input interpretation may be different. Implementations do not normally need to do anything different, except for mappers of the type needed for hidden multicomponent models such as "bym2", which can be handled by <code>bm_collect</code> .
multi	logical; If TRUE (or positive), recurse one level into sub-mappers
input	Data input for the mapper.
state	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>

### Methods (by generic)

- `ibm_eval(bm_taylor)`: Evaluates linearised mapper information at the given state. The `input` argument is ignored, so that the usual argument order `ibm_eval(mapper, input, state)` syntax can be used, but also `ibm_eval(mapper, state = state)`. For a mapper with a named jacobian list, the `state` argument must also be a named list. If `state` is `NULL`, all-zero is assumed.

### See Also

`bru_mapper`, `bru_mapper_generics`

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesh()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_multi()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

### Examples

```
m <- bm_taylor(
  offset = rep(2, 3),
  jacobian = matrix(1:6, 3, 2),
  state0 = c(1, 2)
)
ibm_eval2(m, state = 2:3)
```

## Description

This method is a wrapper for `INLA::inla` and provides multiple enhancements.

- Easy usage of spatial covariates and automatic construction of `inla` projection matrices for (spatial) SPDE models. This feature is accessible via the `components` parameter. Practical examples on how to use spatial data by means of the `components` parameter can also be found by looking at the `lgcp` function's documentation.
- Constructing multiple likelihoods is straight forward. See `like` for more information on how to provide additional likelihoods to `bru` using the `...` parameter list.
- Support for non-linear predictors. See example below.
- Log Gaussian Cox process (LGCP) inference is available by using the `cp` family or (even easier) by using the `lgcp` function.

## Usage

```
bru(components = ~Intercept(1), ..., options = list(), .envir = parent.frame())

bru_rerun(result, options = list())

## S3 method for class 'bru'
print(x, ...)
```

## Arguments

<code>components</code>	A formula-like specification of latent components. Also used to define a default linear additive predictor. See <code>bru_comp()</code> for details.
<code>...</code>	Observation models, each constructed by a calling <code>bru_obs()</code> , or named parameters that can be passed to a single <code>bru_obs()</code> call. Note that all the arguments will be evaluated before calling <code>bru_obs()</code> in order to detect if they are like objects. This means that special arguments that need to be evaluated in the context of <code>response_data</code> or <code>data</code> (such as <code>Ntrials</code> ) may will only work that way in direct calls to <code>bru_obs()</code> .
<code>options</code>	A <code>bru_options</code> options object or a list of options passed on to <code>bru_options()</code>
<code>.envir</code>	Environment for component evaluation (for when a non-formula specification is used)
<code>result</code>	A previous estimation object of class <code>bru</code>
<code>x</code>	A <code>bru</code> object to be printed

## Value

`bru` returns an object of class "bru". A `bru` object inherits from `INLA::inla` (see the `inla` documentation for its properties) and adds additional information stored in the `bru_info` field.

## Methods (by generic)

- `print(bru)`: Print a summary of a `bru` object.

## Functions

- `bru_rerun()`: Continue the optimisation from a previously computed estimate. The estimation options list can be given new values to override the original settings.  
To rerun with a subset of the data (e.g. for cross validation or prior sampling), use `bru_set_missing()` to set all or part of the response data to NA before calling `bru_rerun()`.

## Author(s)

Fabian E. Bachl <bachlfab@gmail.com>

## Examples

```
if (bru_safe_inla()) {
  # Simulate some covariates x and observations y
  input.df <- data.frame(x = cos(1:10))
  input.df <- within(input.df, {
    y <- 5 + 2 * x + rnorm(10, mean = 0, sd = 0.1)
  })

  # Fit a Gaussian likelihood model
  fit <- bru(y ~ x + Intercept(1), family = "gaussian", data = input.df)

  # Obtain summary
  fit$summary.fixed
}

if (bru_safe_inla()) {
  # Alternatively, we can use the bru_obs() function to construct the likelihood:

  lik <- bru_obs(family = "gaussian",
                  formula = y ~ x + Intercept,
                  data = input.df)
  fit <- bru(~ x + Intercept(1), lik)
  fit$summary.fixed
}

# An important addition to the INLA methodology is bru's ability to use
# non-linear predictors. Such a predictor can be formulated via bru_obs()'s
# \code{formula} parameter. The z(1) notation is needed to ensure that
# the z component should be interpreted as single latent variable and not
# a covariate:

if (bru_safe_inla()) {
  z <- 2
  input.df <- within(input.df, {
    y <- 5 + exp(z) * x + rnorm(10, mean = 0, sd = 0.1)
  })
  lik <- bru_obs(
    family = "gaussian", data = input.df,
    formula = y ~ exp(z) * x + Intercept
  )
}
```

```

fit <- bru(~ z(1) + Intercept(1), lik)

# Check the result (z posterior should be around 2)
fit$summary.fixed
}

```

## Description

Similar to `glm()`, `gam()` and `inla()`, `bru()` models can be constructed via a formula-like syntax, where each latent effect is specified. However, in addition to the parts of the syntax compatible with `INLA::inla`, `bru` components offer additional functionality which facilitates modelling, and the predictor expression can be specified separately, allowing more complex and non-linear predictors to be defined. The formula syntax is just a way to allow all model components to be defined in a single line of code, but the definitions can optionally be split up into separate component definitions. See Details for more information.

The `bru_comp` methods all rely on the `bru_comp.character()` method, that defines a model component with a given label/name. The user usually doesn't need to call these methods directly, but can instead supply a formula expression that can be interpreted by the `bru_comp_list.formula()` method, called inside `bru()`.

## Usage

```

bru_comp(...)

bru_component(...)

## S3 method for class 'character'
bru_comp(
  object,
  main = NULL,
  weights = NULL,
  ...,
  model = NULL,
  mapper = NULL,
  main_layer = NULL,
  main_selector = NULL,
  n = NULL,
  values = NULL,
  season.length = NULL,
  nrow = NULL,
  ncol = NULL,
  copy = NULL,
  weights_layer = NULL,

```

```

weights_selector = NULL,
group = 1L,
group_mapper = NULL,
group_layer = NULL,
group_selector = NULL,
ngroup = NULL,
control.group = NULL,
replicate = 1L,
replicate_mapper = NULL,
replicate_layer = NULL,
replicate_selector = NULL,
nrep = NULL,
marginal = NULL,
A.msk = deprecated(),
.envir = parent.frame(),
envir_extra = NULL
)

```

## Arguments

...	Parameters passed on to other methods
object	A character label for the component
main	main takes an R expression that evaluates to where the latent variables should be evaluated (coordinates, indices, continuous scalar (for rw2 etc)). Arguments starting with weights, group, replicate behave similarly to main, but for the corresponding features of INLA::f().
weights, weights_layer, weights_selector	Optional specification of effect scaling weights. Same syntax as for main.
model	Either one of "const" (same as "offset"), "factor_full", "factor_contrast", "linear", "fixed", or a model name or object accepted by INLA's f function. If set to NULL, then "linear" is used for vector inputs, and "fixed" for matrix input (converted internally to an iid model with fixed precision)
mapper	Information about how to do the mapping from the values evaluated in main, and to the latent variables. Auto-detects spde model objects in model and extracts the mesh object to use as the mapper, and auto-generates mappers for indexed models. (Default: NULL, for auto-determination)
main_layer, main_selector	The _layer input should evaluate to a numeric index or character name or vector of which layer/variable to extract from a covariate data object given in main. (Default: NULL if _selector is given. Otherwise the effect component name, if it exists in the covariate object, and otherwise the first column of the covariate data frame) The _selector value should be a character name of a variable whose contents determines which layer to extract from a covariate for each data point. (Default: NULL)
n	The number of latent variables in the model. Should be auto-detected for most or all models. Default: NULL, for auto-detection. Models with matrix input for

	Cmatrix or graph will use the matrix size. An error is given if it realises it can't figure it out by itself.
values	Specifies for what covariate/index values INLA should build the latent model. Normally generated internally based on the mapping details. (Default: NULL, for auto-determination)
season.length	Passed on to INLA::f() for model "seasonal" (TODO: check if this parameter is still fully handled)
nrow, ncol	Number of rows and columns for model types "rw2d", "rw2diid", and "matern2d". Default is NULL.
copy	character; label of other component that this component should be a copy of. If the fixed = FALSE, a scaling constant is estimated, via a hyperparameter. If fixed = TRUE, the component scaling is fixed, by default to 1; for fixed scaling, it's more efficient to express the scaling in the predictor expression instead of making a copy component.
group, group_mapper, group_layer, group_selector, ngroup	Optional specification of kronecker/group model indexing.
control.group	list of kronecker/group model parameters, currently passed directly on to INLA::f replicate, replicate_mapper, replicate_layer, replicate_selector, nrep
	Optional specification of indices for an independent replication model. Same syntax as for main
marginal	May specify a bm_marginal() mapper, that is applied before scaling by weights.
A.msk	[Deprecated] and has no effect.
.envir	Evaluation environment
envir_extra	TODO: check/fix this parameter.

## Details

As shorthand, `bru()` will understand basic additive formulae describing fixed effect models. For instance, the components specification  $y \sim x$  will define the linear combination of an effect named  $x$  and an intercept to the response  $y$  with respect to the likelihood family stated when calling `bru()`. Mathematically, the linear predictor  $\eta$  would be written as

$$\eta = \beta x + c,$$

where:

$c$  is the *intercept*

$x$  is a *covariate*

$\beta$  is a *latent variable* associated with  $x$  and

$\psi = \beta x$  is called the *effect* of  $x$

A problem that arises when using this kind of R formula is that it does not clearly reflect the mathematical formula. For instance, when providing the formula to inla, the resulting object will refer to the random effect  $\psi = \beta * x$  as  $x$ . Hence, it is not clear when  $x$  refers to the covariate or the effect of the covariate.

The `bru_comp.character` method is `inlabru`'s equivalent to INLA's `f()` function but adds functionality that is unique to `inlabru`.

Deprecated parameters:

- map: Use `main` instead.
- mesh: Use `mapper` instead.

## Functions

- `bru_component()`: Backwards compatibility alias for `bru_comp()`

### Naming random effects

In INLA, the `f()` notation is used to define more complex models, but a simple linear effect model can also be expressed as

- `formula = y ~ f(x, model = "linear")`,

where `f()` is the `inla` specific function to set up random effects of all kinds. The underlying predictor would again be  $\eta = \beta * x + c$  but the result of fitting the model would state `x` as the random effect's name. `bru` allows rewriting this formula in order to explicitly state the name of the random effect and the name of the associated covariate. This is achieved by replacing `f` with an arbitrary name that we wish to assign to the effect, e.g.

- `components = y ~ psi(x, model = "linear")`.

Being able to discriminate between  $x$  and  $\psi$  is relevant because of two functionalities `bru` offers. The formula parameters of both `bru()` and the prediction method `predict.bru` are interpreted in the mathematical sense. For instance, `predict` may be used to analyze the analytical combination of the covariate  $x$  and the intercept using

- `predict(fit, data.frame(x=2), ~ exp(psi + Intercept))`.

which corresponds to the mathematical expression  $e^{x\beta+c}$ .

On the other hand, `predict` may be used to only look at a transformation of the latent variable  $\beta_\psi$

- `predict(fit, NULL, ~ exp(psi_latent))`.

which corresponds to the mathematical expression  $e^\beta$ .

### Author(s)

Fabian E. Bachl <[bachlfab@gmail.com](mailto:bachlfab@gmail.com)> and Finn Lindgren <[Finn.Lindgren@gmail.com](mailto:Finn.Lindgren@gmail.com)>

### See Also

[bru\\_input\(\)](#), [summary.bru\\_comp\(\)](#)

Other component constructors: [bru\\_comp\\_list\(\)](#)

## Examples

```

# As an example, let us create a linear component. Here, the component is
# called "myLinearEffectOfX" while the covariate the component acts on is
# called "x". Note that a list of components is returned because the
# formula may define multiple components

cmp <- bru_comp_list(~ myLinearEffectOfX(main = x, model = "linear"))
summary(cmp)
# Equivalent shortcuts:
cmp <- bru_comp_list(~ myLinearEffectOfX(x, model = "linear"))
cmp <- bru_comp_list(~ myLinearEffectOfX(x))
# Individual component
cmp <- bru_comp("myLinearEffectOfX", main = x, model = "linear")
summary(cmp)

if (bru_safe_inla()) {
  # As an example, let us create a linear component. Here, the component is
  # called "myEffectOfX" while the covariate the component acts on is called
  # "x":

  cmp <- bru_comp("myEffectOfX", main = x, model = "linear")
  summary(cmp)

  # A more complicated component:
  cmp <- bru_comp("myEffectOfX",
    main = x,
    model = INLA::inla.spde2.matern(fm_mesh_1d(1:10))
  )

  # Compound fixed effect component, where x and z are in the input data.
  # The formula will be passed on to MatrixModels::model.Matrix:
  cmp <- bru_comp("eff", ~ -1 + x:z, model = "fixed")
  summary(cmp)
}

```

**bru\_comp\_eval**

*Evaluate component values in predictor expressions*

## Description

In predictor expressions, `name_eval(...)` can be used to evaluate the effect of a component called "name".

## Usage

```
bru_comp_eval(
  main,
```

```

group = NULL,
replicate = NULL,
weights = NULL,
.state = NULL
)

```

### Arguments

`main, group, replicate, weights`

Specification of where to evaluate a component. The four inputs are passed on to the joint `bru_mapper` for the component, as

```

list(mapper = list(
  main = main,
  group = group,
  replicate = replicate),
  scale = weights)

```

NOTE: If you have model component with the same name as a data variable you want to supply as input to `name_eval()`, you need to use `.data.[["myvar"]]` to access it. Otherwise, it will try to use the other component effect as input, which is ill-defined.

`.state`

The internal component state. Normally supplied automatically by the internal methods for evaluating inlabru predictor expressions.

### Value

A vector of values for a component

### Examples

```

if (bru_safe_inla() &&
  require("sf", quietly = TRUE) &&
  requireNamespace("sn", quietly = TRUE)) {
  mesh <- fmesher::fm_mesh_2d_inla(
    cbind(0, 0),
    offset = 2,
    max.edge = 2.5
  )
  spde <- INLA::inla.spde2.pcmatern(
    mesh,
    prior.range = c(1, NA),
    prior.sigma = c(0.2, NA)
  )
  set.seed(12345L)
  data <- sf::st_as_sf(
    data.frame(
      x = runif(50),
      y = runif(50),
      z = rnorm(50)
    ),
    
```

```

    coords = c("x", "y")
  )
  fit <- bru(
    z ~ -1 + field(geometry, model = spde),
    family = "gaussian", data = data,
    options = list(control.inla = list(int.strategy = "eb")))
  )
  pred <- generate(
    fit,
    newdata = data.frame(A = 0.5, B = 0.5),
    formula = ~ field_eval(cbind(A, B)),
    n.samples = 1L
  )
}

```

**bru\_comp\_list***Methods for inlabru component lists***Description**

Constructor methods for inlabru component lists. Syntax details are given in [bru\\_comp\(\)](#).

**Usage**

```

bru_comp_list(object, lhoods = NULL, .envir = parent.frame(), ...)

## S3 method for class 'formula'
bru_comp_list(object, lhoods = NULL, .envir = parent.frame(), ...)

## S3 method for class 'list'
bru_comp_list(
  object,
  lhoods = NULL,
  .envir = parent.frame(),
  inputs = NULL,
  ...
)

## S3 method for class 'bru_comp_list'
c(...)

## S3 method for class 'bru_comp'
c(...)

## S3 method for class 'bru_comp_list'
x[i]

```

## Arguments

<code>object</code>	The object to operate on
<code>lhoods</code>	A <a href="#">bru_obs_list</a> object
<code>.envir</code>	An evaluation environment for non-formula input
<code>...</code>	Parameters passed on to other methods. Also see Details.
<code>inputs</code>	A tree-like list of component input evaluations, from <a href="#">bru_input.bru_obs_list()</a> .
<code>x</code>	<code>bru_comp_list</code> object from which to extract a sub-list
<code>i</code>	indices specifying elements to extract

## Methods (by class)

- `bru_comp_list(formula)`: Convert a component formula into a `bru_comp_list` object
- `bru_comp_list(list)`: Combine a list of components and/or component formulas into a `bru_comp_list` object

## Methods (by generic)

- `c(bru_comp_list)`: The ... arguments should be `bru_comp_list` objects. The environment from the first argument will be applied to the resulting `bru_comp_list`.

## Functions

- `c(bru_comp)`: The ... arguments should be component objects from [bru\\_comp\(\)](#). The environment from the first argument will be applied to the resulting `bru_comp_list`.

## Author(s)

Fabian E. Bachl <bachlfab@gmail.com> and Finn Lindgren <finn.lindgren@gmail.com>

## See Also

Other component constructors: [bru\\_comp\(\)](#)  
 Other component constructors: [bru\\_comp\(\)](#)

## Examples

```
# As an example, let us create a linear component. Here, the component is
# called "myLinearEffectOfX" while the covariate the component acts on is
# called "x". Note that a list of components is returned because the
# formula may define multiple components

eff <- bru_comp_list(~ myLinearEffectOfX(main = x, model = "linear"))
summary(eff[[1]])
# Equivalent shortcuts:
eff <- bru_comp_list(~ myLinearEffectOfX(x, model = "linear"))
eff <- bru_comp_list(~ myLinearEffectOfX(x))
# Individual component
eff <- bru_comp("myLinearEffectOfX", main = x, model = "linear")
```

---

**bru\_convergence\_plot** *Plot inlabru convergence diagnostics*

---

## Description

Draws four panels of convergence diagnostics for an iterated INLA method estimation

## Usage

```
bru_convergence_plot(x, from = 1, to = NULL, type = NULL)
```

## Arguments

x	a <b>bru</b> object, typically a result from <b>bru()</b> for a nonlinear predictor model
from, to	integer values for the range of iterations to plot. Default <b>from</b> = 1 (start from the first iteration) and <b>to</b> = NULL (end at the last iteration). Set <b>from</b> = 0 to include the initial linearisation point in the track plot.
type	[Experimental] character; "bru" (default) for iterative nonlinear inlabru convergence diagnostics plots, or "inla" for INLA optimiser trace plots.

## Details

Requires the "dplyr", "ggplot2", "magrittr", and "patchwork" packages to be installed.

## Value

A ggplot object with four panels of convergence diagnostics:

- Tracks: Mode and linearisation values for each effect
- Mode - Lin: Difference between mode and linearisation values for each effect
- |Change| / sd: Absolute change in mode and linearisation values divided by the standard deviation for each effect
- Change & sd: Absolute change in mode and linearisation values and standard deviation for each effect

For multidimensional components, only the overall average, maximum, and minimum values are shown.

## See Also

[bru\(\)](#)

## Examples

```
## Not run:  
fit <- bru(...)  
bru_convergence_plot(fit)  
  
## End(Not run)
```

---

bru_fill_missing	<i>Fill in missing values in Spatial grids</i>
------------------	--

---

## Description

Computes nearest-available-value imputation for missing values in space

## Usage

```
bru_fill_missing(
  data,
  where,
  values,
  layer = NULL,
  selector = NULL,
  batch_size = deprecated()
)
```

## Arguments

<code>data</code>	A <code>SpatialPointsDataFrame</code> , <code>SpatialPixelsDataFrame</code> , <code>SpatialGridDataFrame</code> , <code>Spa-</code> <code>tRaster</code> , <code>Raster</code> , or <code>sf</code> object containing data to use for filling
<code>where</code>	<code>A</code> , <code>matrix</code> , <code>data.frame</code> , or <code>SpatialPoints</code> or <code>SpatialPointsDataFrame</code> , or <code>sf</code> object, containing the locations of the evaluated values
<code>values</code>	A vector of values to be filled in where <code>is.na(values)</code> is <code>TRUE</code>
<code>layer</code> , <code>selector</code>	Specifies what data column or columns from which to extract data, see <code>bru_comp()</code> for details.
<code>batch_size</code>	<b>[Deprecated]</b> due to improved algorithm. Size of nearest-neighbour calculation blocks, to limit the memory and computational complexity.

## Value

An infilled vector of values

## Examples

```
## Not run:
if (bru_safe_inla()) {
  points <-
    sp::SpatialPointsDataFrame(
      matrix(1:6, 3, 2),
      data = data.frame(val = c(NA, NA, NA))
    )
  input_coord <- expand.grid(x = 0:7, y = 0:7)
  input <-
    sp::SpatialPixelsDataFrame(
      input_coord,
```

```

        data = data.frame(val = as.vector(input_coord$y))
    )
points$val <- bru_fill_missing(input, points, points$val)
print(points)

# To fill in missing values in a grid:
print(input$val[c(3, 30)])
input$val[c(3, 30)] <- NA # Introduce missing values
input$val <- bru_fill_missing(input, input, input$val)
print(input$val[c(3, 30)])
}

## End(Not run)

```

**bru\_get\_mapper***Extract mapper information from INLA model component objects***Description**

The component definitions will automatically attempt to extract mapper information from any model object by calling the generic `bru_get_mapper`. Any class method implementation should return a `bru_mapper` object suitable for the given latent model.

**Usage**

```

bru_get_mapper(model, ...)

## S3 method for class 'inla.spde'
bru_get_mapper(model, ...)

## S3 method for class 'inla.rgeneric'
bru_get_mapper(model, ...)

bru_get_mapper_safely(model, ...)

```

**Arguments**

<code>model</code>	A model component object
<code>...</code>	Arguments passed on to other methods

**Value**

A `bru_mapper` object defined by the model component

### Methods (by class)

- **`bru_get_mapper(inla.spde)`**: Extract an indexed mapper for the `model$mesh` object contained in the `model` object, which is assumed to be of a class supporting relevant `fmesher` methods.
- **`bru_get_mapper(inla.rgeneric)`**: Returns the mapper given by a call to `model$f$rgeneric$definition("mapper")`. To support this for your own `inla.rgeneric` models, add a "mapper" option to the `cmd` argument of your `rgeneric` definition function. You will need to store the mapper in your object as well. Alternative, define your model using a subclass and define a corresponding `bru_get_mapper.subclass` method that should return the corresponding `bru_mapper` object.

### Functions

- **`bru_get_mapper_safely()`**: Tries to call the `bru_get_mapper`, and returns `NULL` if it fails (e.g. due to no available class method). If the call succeeds and returns non-`NULL`, it checks that the object inherits from the `bru_mapper` class, and gives an error if it does not.

### See Also

`bru_mapper` for mapper constructor methods, and the individual mappers for specific implementation details.

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesher()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_multi()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bm_taylor()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

### Examples

```
if (bru_safe_inla()) {
  library(INLA)
  mesh <- fmesher::fm_rcdt_2d_inla(globe = 2)
  spde <- inla.spde2.pcmatern(mesh,
    prior.range = c(1, 0.5),
    prior.sigma = c(1, 0.5)
  )
  mapper <- bru_get_mapper(spde)
  ibm_n(mapper)
}
```

### Description

The `bru_info` class is used to store metadata about `bru` models.

**Usage**

```

bru_info(...)

## S3 method for class 'character'
bru_info(method, ..., inlabru_version = NULL, INLA_version = NULL)

## S3 method for class 'bru'
bru_info(object, ...)

## S3 method for class 'bru_info'
summary(object, verbose = TRUE, ...)

## S3 method for class 'summary_bru_info'
print(x, ...)

## S3 method for class 'bru_info'
print(x, ...)

```

**Arguments**

...	Arguments passed on to other <code>summary</code> methods
method	character; The type of estimation method used
inlabru_version	character; inlabru package version. Default: NULL, for automatically detecting the version
INLA_version	character; INLA package version. Default: NULL, for automatically detecting the version
object	Object to operate on
verbose	logical; If TRUE, include more details of the component definitions. If FALSE, only show basic component definition information. Default: TRUE
x	An object to be printed

**Methods (by class)**

- `bru_info(character)`: Create a `bru_info` object
- `bru_info(bru)`: Extract the `bru_info` object from an estimated `bru()` result object. The default print method show information about model components and observation models.

## Description

Access method for *bru\_log* objects. Note: Up to version 2.8.0, *bru\_log()* was a deprecated alias for *bru\_log\_message()*. When running on 2.8.0 or earlier, use *bru\_log\_get()* to access the global log, and *cat(fit\$bru\_iinla\$log, sep = "\n")* to print a stored estimation object log. After version 2.8.0, use *bru\_log()* to access the global log, and *bru\_log(fit)* to access a stored estimation log.

## Usage

```
bru_log(x = NULL, verbosity = NULL)

## S3 method for class 'character'
bru_log(x, verbosity = NULL)

## S3 method for class 'bru_log'
bru_log(x, verbosity = NULL)

## S3 method for class 'iinla'
bru_log(x, verbosity = NULL)

## S3 method for class 'bru'
bru_log(x, verbosity = NULL)

## S3 method for class 'bru_log'
format(x, ..., timestamp = TRUE, verbosity = FALSE)

## S3 method for class 'bru_log'
print(x, ..., timestamp = TRUE, verbosity = FALSE)

## S3 method for class 'bru_log'
as.character(x, ...)

## S3 method for class 'bru_log'
x[i]

## S3 method for class 'bru_log'
c(...)

## S3 method for class 'bru_log'
length(x)
```

## Arguments

<i>x</i>	An object that is, contains, or can be converted to, a <i>bru_log</i> object. If NULL, refers to the global <i>iinlabru</i> log.
<i>verbosity</i>	integer value for limiting the highest verbosity level being returned.
...	further arguments passed to or from other methods.
<i>timestamp</i>	If TRUE, include the timestamp of each message. Default TRUE.

- i indices specifying elements to extract. If character, denotes the sequence between bookmark i and the next bookmark (or the end of the log if i is the last bookmark)

## Value

`bru_log` A `bru_log` object, containing a character vector of log messages, and potentially a vector of bookmarks.

## Methods (by generic)

- `format(bru_log)`: Format a `bru_log` object for printing. If `verbosity` is TRUE, include the `verbosity` level of each message.
- `print(bru_log)`: Print a `bru_log` object with `cat(x, sep = "\n")`. If `verbosity` is TRUE, include the `verbosity` level of each message.
- `as.character(bru_log)`: Convert `bru_log` object to a plain character vector
- `[: Extract a subset of a bru_log object`
- `c(bru_log)`: Concatenate several `bru_log` or `character` objects into a `bru_log` object.
- `length(bru_log)`: Obtain the number of log entries into a `bru_log` object.

## Functions

- `bru_log()`: Extract stored log messages. If non-NULL, the `verbosity` argument determines the maximum `verbosity` level of the messages to extract.

## See Also

Other inlabru log methods: `bru_log_bookmark()`, `bru_log_message()`, `bru_log_new()`, `bru_log_offset()`, `bru_log_reset()`

## Examples

```
bru_log(verbosity = 2L)
format(bru_log())

bru_log(verbosity = 2L)
print(bru_log(), timestamp = TRUE, verbosity = TRUE)
```

---

**bru\_log\_bookmark**      *Methods for bru\_log bookmarks*

---

**Description**

Methods for `bru_log` bookmarks.

**Usage**

```
bru_log_bookmark(bookmark = "", offset = NULL, x = NULL)  
bru_log_bookmarks(x = NULL)
```

**Arguments**

<code>bookmark</code>	character; The label for a bookmark with a stored offset.
<code>offset</code>	integer; a position offset in the log, with <code>0L</code> pointing at the start of the log. If negative, denotes the point <code>abs(offset)</code> elements from tail of the log. When <code>bookmark</code> is non-NULL, the <code>offset</code> applies a shift (forwards or backwards) to the bookmark list.
<code>x</code>	A <code>bru_log</code> object. If <code>NULL</code> , the global <code>inlabru</code> log is used.

**Value**

`bru_log_bookmark()`: Returns the modified `bru_log` object if `x` is non-NULL.  
`bru_log_bookmarks()`: Returns the bookmark vector associated with `x`

**Functions**

- `bru_log_bookmark()`: Set a log bookmark. If `offset` is `NULL` (the default), the bookmark will point to the current end of the log.
- `bru_log_bookmarks()`: Return a integer vector with named elements being bookmarks into the global `inlabru` log with associated log position offsets.

**See Also**

Other `inlabru` log methods: [bru\\_log\(\)](#), [bru\\_log\\_message\(\)](#), [bru\\_log\\_new\(\)](#), [bru\\_log\\_offset\(\)](#), [bru\\_log\\_reset\(\)](#)

---

bru\_log\_message      *Add a log message*

---

## Description

Adds a log message.

## Usage

```
bru_log_message(  
  ...,  
  domain = NULL,  
  appendLF = TRUE,  
  verbosity = 1L,  
  allow_verbose = TRUE,  
  verbose = NULL,  
  verbose_store = NULL,  
  x = NULL  
)  
  
bru_log_abort(  
  msg,  
  ...,  
  domain = NULL,  
  appendLF = TRUE,  
  verbosity = 1L,  
  allow_verbose = TRUE,  
  verbose = FALSE,  
  verbose_store = NULL,  
  call = rlang::caller_env(),  
  .frame = rlang::caller_env()  
)  
  
bru_log_warn(  
  msg,  
  ...,  
  domain = NULL,  
  appendLF = TRUE,  
  verbosity = 1L,  
  allow_verbose = TRUE,  
  verbose = FALSE,  
  verbose_store = NULL,  
  call = rlang::caller_env(),  
  .frame = rlang::caller_env()  
)
```

## Arguments

...	For <code>bru_log_message()</code> , zero or more objects passed on to <code>base:::makeMessage()</code> . For <code>bru_log_abort()</code> and <code>bru_log_warn()</code> , passed on to <code>rlang:::abort()</code> and <code>rlang:::warn()</code> .
domain	Domain for translations, passed on to <code>base:::makeMessage()</code>
appendLF	logical; whether to add a newline to the message. Only used for verbose output.
verbosity	numeric value describing the verbosity level of the message
allow_verbose	Whether to allow verbose output. Must be set to FALSE until the options object has been initialised.
verbose	logical, numeric, or NULL; local override for verbose output. If NULL, the global option <code>bru_verbose</code> or default value is used. If FALSE, no messages are printed. If TRUE, messages with <code>verbosity ≤ 1</code> are printed. If numeric, messages with <code>verbosity ≤ verbose</code> are printed.
verbose_store	Same as <code>verbose</code> , but controlling what messages are stored in the global log object. Can be controlled via the <code>bru_verbose_store</code> with <code>bru_options_set()</code> .
x	A <code>bru_log</code> object. If NULL, refers to the global <code>inlabru</code> log.
msg	character; passed to <code>base:::makeMessage()</code>
call	The calling environment.
.frame	The throwing context, for when <code>.internal</code> is TRUE

## Value

`bru_log_message` returns `invisible(x)`, where `x` is the updated `bru_log` object, or NULL.

## Functions

- `bru_log_abort()`: Store a log message and throw an error.
- `bru_log_warn()`: Store a log message and throw a warning.

## See Also

Other `inlabru` log methods: `bru_log()`, `bru_log_bookmark()`, `bru_log_new()`, `bru_log_offset()`, `bru_log_reset()`

## Examples

```
if (interactive()) {
  code_runner <- function() {
    bru_options_set_local(
      # Show messages up to and including level 2 (default 0)
      bru_verbose = 2,
      # Store messages to an including level 3 (default Inf, storing all)
      bru_verbose_store = 3
    )
    bru_log_bookmark("bookmark 1")
  }
}
```

```
    bru_log_message("Test message 1", verbosity = 1)
    bru_log_message("Test message 2", verbosity = 2)
    bru_log_bookmark("bookmark 2")
    bru_log_message("Test message 3", verbosity = 3)
    bru_log_message("Test message 4", verbosity = 4)

    invisible()
}
message("Run code")
code_runner()
message("Check log from bookmark 1")
print(bru_log()["bookmark 1"])
message("Check log from bookmark 2")
print(bru_log()["bookmark 2"])
}
```

---

**bru\_log\_new** *Create a bru\_log object*

---

## Description

Create a bru\_log object, by default empty.

## Usage

```
bru_log_new(x = NULL, bookmarks = NULL)
```

## Arguments

- |           |   |
|-----------|---|
| x         | An optional character vector of log messages, or data.frame with columns message, timestamp, and verbosity. |
| bookmarks | An optional integer vector of named bookmarks message in x.   |

## See Also

Other inlabru log methods: [bru\\_log\(\)](#), [bru\\_log\\_bookmark\(\)](#), [bru\\_log\\_message\(\)](#), [bru\\_log\\_offset\(\)](#), [bru\\_log\\_reset\(\)](#)

## Examples

```
x <- bru_log_new()
x <- bru_log_message("Test message", x = x)
print(x)
```

---

<b><i>bru_log_offset</i></b>	<i>Position methods for bru_log objects</i>
------------------------------	---

---

## Description

Position methods for **bru\_log** objects.

## Usage

```
bru_log_offset(x = NULL, bookmark = NULL, offset = NULL)

bru_log_index(x = NULL, i, verbosity = NULL)
```

## Arguments

<b>x</b>	A <b>bru_log</b> object. If <b>NULL</b> , the global <b>inlabru</b> log is used.
<b>bookmark</b>	character; The label for a bookmark with a stored offset.
<b>offset</b>	integer; a position offset in the log, with <b>0L</b> pointing at the start of the log. If negative, denotes the point <b>abs(offset)</b> elements from tail of the log. When <b>bookmark</b> is non- <b>NULL</b> , the <b>offset</b> applies a shift (forwards or backwards) to the bookmark list.
<b>i</b>	indices specifying elements to extract. If <b>character</b> , denotes the sequence between bookmark <b>i</b> and the next bookmark (or the end of the log if <b>i</b> is the last bookmark)
<b>verbosity</b>	integer value for limiting the highest verbosity level being returned.

## Functions

- **bru\_log\_offset()**: Utility function for computing log position offsets.
- **bru\_log\_index()**: Utility function for computing index vectors for **bru\_log** objects.

## See Also

Other **inlabru** log methods: [bru\\_log\(\)](#), [bru\\_log\\_bookmark\(\)](#), [bru\\_log\\_message\(\)](#), [bru\\_log\\_new\(\)](#), [bru\\_log\\_reset\(\)](#)

---

bru_log_reset	<i>Clear log contents</i>
---------------	---------------------------

---

## Description

Clears the log contents up to a given offset or bookmark. Default: clear the entire log. When `x` is `NULL`, the global `inlabru` log is updated, and `invisible(NULL)` is returned. Otherwise the updated object is returned (invisibly).

## Usage

```
bru_log_reset(x = NULL, bookmark = NULL, offset = NULL)
```

## Arguments

- |                       |   |
|-----------------------|---|
| <code>x</code>        | A <code>bru_log</code> object, or in some cases, and object that can be converted/extracted to a <code>bru_log</code> object. <code>NULL</code> denotes the global <code>inlabru</code> log object.   |
| <code>bookmark</code> | character; The label for a bookmark with a stored offset.   |
| <code>offset</code>   | integer; a position offset in the log, with <code>0L</code> pointing at the start of the log. If negative, denotes the point <code>abs(offset)</code> elements from tail of the log. When <code>bookmark</code> is non- <code>NULL</code> , the <code>offset</code> applies a shift (forwards or backwards) to the bookmark list. |

## Value

Returns (invisibly) the modified `bru_log` object, or `NULL` (when `x` is `NULL`)

## See Also

Other `inlabru` log methods: [bru\\_log\(\)](#), [bru\\_log\\_bookmark\(\)](#), [bru\\_log\\_message\(\)](#), [bru\\_log\\_new\(\)](#), [bru\\_log\\_offset\(\)](#)

## Examples

```
## Not run:  
if (interactive()) {  
  bru_log_reset()  
}  
  
## End(Not run)
```

---

**bru\_mapper***Constructors for bru\_mapper objects*

---

**Description**

Constructors for `bru_mapper` objects

**Usage**

```
bru_mapper(...)
```

```
bru_mapper_define(mapper, new_class = NULL, remove_class = "list", ...)
```

**Arguments**

...	Arguments passed on to sub-methods, or used for special purposes, see details for each function below.
<code>mapper</code>	For <code>bru_mapper_define</code> , a prototype mapper object, see Details.
<code>new_class</code>	If non-NULL, this is added at the front of the class definition
<code>remove_class</code>	If non-NULL, this class or classes is removed from the class definition before adding the <code>new_class</code> names. Default is "list".

**Value**

- `bru_mapper()` returns a `bru_mapper` object

**Functions**

- `bru_mapper()`: Generic mapper S3 constructor, used for constructing mappers for special objects. See below for details of the default constructor `bru_mapper_define()` that can be used to define new mappers in user code.
- `bru_mapper_define()`: Adds the `new_class` and "bru\_mapper" class names to the inheritance list for the input `mapper` object, unless the object already inherits from these.

To register mapper classes and methods in scripts, use `.S3method()` to register the methods, e.g. `.S3method("ibm_jacobian", "my_mapper_class", ibm_jacobian.my_mapper_class)`.

In packages with `Suggests: inlabru`, add method information for delayed registration, e.g.:

```
#' @rawNamespace S3method(inlabru::bru_get_mapper, inla_rspde)
#' @rawNamespace S3method(inlabru::ibm_n, bru_mapper_inla_rspde)
#' @rawNamespace S3method(inlabru::ibm_values, bru_mapper_inla_rspde)
#' @rawNamespace S3method(inlabru::ibm_jacobian, bru_mapper_inla_rspde)
```

or before each method, use `@exportS3Method`:

```
#' @exportS3Method inlabru::bru_get_mapper
```

etc., which semi-automates it.

## See Also

[bru\\_mapper\\_generics](#) for generic methods, the individual mapper pages for special method implementations, and [bru\\_get\\_mapper](#) for hooks to extract mappers from latent model object class objects.

Other mappers: [bm\\_aggregate\(\)](#), [bm\\_collect\(\)](#), [bm\\_const\(\)](#), [bm\\_factor\(\)](#), [bm\\_fmesh\(\)](#), [bm\\_harmonics\(\)](#), [bm\\_index\(\)](#), [bm\\_linear\(\)](#), [bm\\_logsumexp\(\)](#), [bm\\_marginal\(\)](#), [bm\\_matrix\(\)](#), [bm\\_mesh\\_B\(\)](#), [bm\\_multi\(\)](#), [bm\\_pipe\(\)](#), [bm\\_repeat\(\)](#), [bm\\_scale\(\)](#), [bm\\_shift\(\)](#), [bm\\_sum\(\)](#), [bm\\_taylor\(\)](#), [bru\\_get\\_mapper\(\)](#), [bru\\_mapper.fm\\_mesh\\_1d\(\)](#), [bru\\_mapper.fm\\_mesh\\_2d\(\)](#), [bru\\_mapper\\_generics](#)

## Examples

```
mapper <- bm_index(5)
ibm_jacobian(mapper, input = c(1, 3, 4, 5, 2))
```

**bru\_mapper.fm\_mesh\_1d** *Mapper for fm\_mesh\_1d*

## Description

Create mapper for an fm\_mesh\_1d object

## Usage

```
## S3 method for class 'fm_mesh_1d'
bru_mapper(mesh, indexed = TRUE, ...)

## S3 method for class 'bm_fm_mesh_1d'
ibm_n(mapper, ...)

## S3 method for class 'bm_fm_mesh_1d'
ibm_values(mapper, ...)

## S3 method for class 'bm_fm_mesh_1d'
ibm_jacobian(mapper, input, ...)

## S3 method for class 'bm_inla_mesh_1d'
ibm_n(mapper, ...)

## S3 method for class 'bm_inla_mesh_1d'
ibm_values(mapper, ...)

## S3 method for class 'bm_inla_mesh_1d'
ibm_jacobian(mapper, input, ...)
```

### Arguments

<code>mesh</code>	An <code>fm_mesh_1d</code> object to use as a mapper
<code>indexed</code>	logical; If TRUE (default), the <code>ibm_values()</code> output will be the integer indexing sequence for the latent variables (needed for spde models). If FALSE, points representative of the basis centres are returned (useful for an interpolator for rw2 models and similar, for fmesher versions >= 0.3.0.9002).
<code>...</code>	Arguments passed on to other methods
<code>mapper</code>	A mapper S3 object, inheriting from <code>bru_mapper</code> .
<code>input</code>	Data input for the mapper.

### Value

A `bm_fm_mesh_1d` or `bm_fmesher` object. The the general `bm_fmesher()` mapper handles all indexed fmesher objects.

### See Also

[bru\\_mapper](#), [bru\\_mapper\\_generics](#)

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesher()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_multi()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_2d()`, `bru_mapper_generics`

### Examples

```
m <- bru_mapper(fm_mesh_1d(c(1:3, 5, 7)))
ibm_values(m)
ibm_eval(m, 1:7, 1:5)

m <- bru_mapper(fm_mesh_1d(c(1:3, 5, 7)), indexed = FALSE)
ibm_values(m)
ibm_eval(m, 1:7, 1:5)

m <- bru_mapper(
  fm_mesh_1d(c(1:3, 5, 7), degree = 2, boundary = "free"),
  indexed = FALSE
)
ibm_values(m)
ibm_eval(m, 1:7, 1:6)
```

`bru_mapper.fm_mesh_2d` *Mapper for fm\_mesh\_2d*

### Description

Creates a mapper for 2D `fm_mesh_2d` objects. Equivalent to calling `bm_fmesher()`.

## Usage

```
## S3 method for class 'fm_mesh_2d'
bru_mapper(mesh, ...)

## S3 method for class 'bm_fm_mesh_2d'
ibm_n(mapper, ...)

## S3 method for class 'bm_fm_mesh_2d'
ibm_values(mapper, ...)

## S3 method for class 'bm_fm_mesh_2d'
ibm_jacobian(mapper, input, ...)

## S3 method for class 'bm_inla_mesh_2d'
ibm_n(mapper, ...)

## S3 method for class 'bm_inla_mesh_2d'
ibm_values(mapper, ...)

## S3 method for class 'bm_inla_mesh_2d'
ibm_jacobian(mapper, input, ...)
```

## Arguments

mesh	An fm_mesh_2d object to use as a mapper
...	Arguments passed on to other methods
mapper	A mapper S3 object, inheriting from bru_mapper.
input	Data input for the mapper.

## Value

A bm\_fmesher object. Note: Prior to version 2.12.0.9021, this was a bru\_mapper\_fm\_mesh\_2d object. Also see the note for [bru\\_mapper.fm\\_mesh\\_1d\(\)](#).

## See Also

[bru\\_mapper](#), [bru\\_mapper\\_generics](#)

Other mappers: [bm\\_aggregate\(\)](#), [bm\\_collect\(\)](#), [bm\\_const\(\)](#), [bm\\_factor\(\)](#), [bm\\_fmesher\(\)](#), [bm\\_harmonics\(\)](#), [bm\\_index\(\)](#), [bm\\_linear\(\)](#), [bm\\_logsumexp\(\)](#), [bm\\_marginal\(\)](#), [bm\\_matrix\(\)](#), [bm\\_mesh\\_B\(\)](#), [bm\\_multi\(\)](#), [bm\\_pipe\(\)](#), [bm\\_repeat\(\)](#), [bm\\_scale\(\)](#), [bm\\_shift\(\)](#), [bm\\_sum\(\)](#), [bm\\_taylor\(\)](#), [bru\\_get\\_mapper\(\)](#), [bru\\_mapper\(\)](#), [bru\\_mapper.fm\\_mesh\\_1d\(\)](#), [bru\\_mapper\\_generics](#)

## Examples

```
m <- bru_mapper(fmesher::fmexample$mesh)
ibm_n(m)
ibm_eval(m, as.matrix(expand.grid(-2:2, -2:2)), seq_len(ibm_n(m)))
```

**bru\_mapper\_generics    *Generic methods for bru\_mapper objects***

### Description

A `bru_mapper` sub-class implementation must provide an `ibm_jacobian()` method. If the model size 'n' and definition values 'values' are stored in the object itself, default methods are available (see Details). Otherwise the `ibm_n()` and `ibm_values()` methods also need to be provided.

### Usage

```
ibm_n(mapper, inla_f = FALSE, ...)

ibm_n_output(mapper, input, state = NULL, inla_f = FALSE, ...)

ibm_values(mapper, inla_f = FALSE, ...)

ibm_is_linear(mapper, ...)

ibm_jacobian(mapper, input, state = NULL, inla_f = FALSE, ...)

ibm_linear(mapper, input, state = NULL, ...)

ibm_simplify(mapper, input = NULL, state = NULL, ...)

ibm_eval(mapper, input, state = NULL, ...)

ibm_eval2(mapper, input, state = NULL, ...)

ibm_names(mapper)

ibm_names(mapper) <- value

ibm_inla_subset(mapper, ...)

ibm_invalid_output(mapper, input, state, ...)

## Default S3 method:
ibm_n(mapper, inla_f = FALSE, ...)

## Default S3 method:
ibm_n_output(mapper, input, state = NULL, inla_f = FALSE, ...)

## Default S3 method:
ibm_values(mapper, inla_f = FALSE, ...)

## Default S3 method:
```

```

ibm_is_linear(mapper, ...)

## Default S3 method:
ibm_jacobian(mapper, input, state = NULL, ...)

## Default S3 method:
ibm_linear(mapper, input, state, ...)

## Default S3 method:
ibm_simplify(mapper, input = NULL, state = NULL, ...)

## Default S3 method:
ibm_eval(mapper, input, state = NULL, ..., jacobian = NULL)

## Default S3 method:
ibm_eval2(mapper, input, state, ...)

## Default S3 method:
ibm_names(mapper, ...)

## Default S3 method:
ibm_inla_subset(mapper, ...)

## Default S3 method:
ibm_invalid_output(mapper, input, state, ...)

```

## Arguments

<code>mapper</code>	A mapper S3 object, inheriting from <code>bru_mapper</code> .
<code>inla_f</code>	logical; when TRUE for <code>ibm_n()</code> and <code>ibm_values()</code> , the result must be compatible with the <code>INLA::f(...)</code> and corresponding <code>INLA::inla.stack(...)</code> constructions. For <code>ibm_{eval,jacobian,linear}</code> , the input interpretation may be different. Implementations do not normally need to do anything different, except for mappers of the type needed for hidden multicomponent models such as "bym2", which can be handled by <code>bm_collect</code> .
<code>...</code>	Arguments passed on to other methods
<code>input</code>	Data input for the mapper.
<code>state</code>	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
<code>value</code>	a character vector of the same length as the number of sub-mappers in the mapper
<code>jacobian</code>	For <code>ibm_eval()</code> methods, an optional pre-computed Jacobian, typically supplied by internal methods that already have the Jacobian.

## Functions

- `ibm_n()`: Implementations must return the size of the latent vector being mapped to.

- `ibm_n_output()`: Implementations must return an integer denoting the mapper output length. The default implementation returns `NROW(input)`. Mappers such as `bm_multi` and `bm_collect`, that can accept `list()` inputs require their own methods implementations.
- `ibm_values()`: When `inla_f=TRUE`, implementations must return a vector that would be interpretable by an `INLA::f(..., values = ...)` specification. The exception is the method for `bm_multi`, that returns a multi-column data frame if `multi=TRUE`.
- `ibm_is_linear()`: Implementations must return TRUE or FALSE. If TRUE (returned by the default method unless the mapper contains an `is_linear` variable), users of the mapper may assume the mapper is linear.
- `ibm_jacobian()`: Implementations must return a (sparse) matrix of size `ibm_n_output(mapper, input, inla_f)` by `ibm_n(mapper, inla_f = FALSE)`. The `inla_f=TRUE` argument should only affect the allowed type of input format.
- `ibm_linear()`: Implementations must return a `bm_taylor` object. The linearisation information includes `offset`, `jacobian`, and `state0`. The state information indicates for which state the offset was evaluated, with `NULL` meaning all-zero. The linearised mapper output is defined as

```
effect(input, state) =
  offset(input, state0) + jacobian(input, state0) %*% (state - state0)
```

The default method calls `ibm_eval()` and `ibm_jacobian()` to generate the needed information.

- `ibm_simplify()`: Implementations must return a `bru_mapper` object. The default method returns `ibm_linear(...)` for linear mappers, and the original mapper for non-linear mappers.
- `ibm_eval()`: Implementations must return a vector of length `ibm_n_output(...)`. The input contents must be in a format accepted by `ibm_jacobian(...)` for the mapper.
- `ibm_eval2()`: Implementations must return a list with elements `offset` and `jacobian`. The input contents must be in a format accepted by `ibm_jacobian(...)` for the mapper.
- `ibm_names()`: Implementations must return a character vector of sub-mapper names, or `NULL`. Intended for providing information about multi-mappers and mapper collections.
- `ibm_names(mapper) <- value`: Set mapper names.
- `ibm_inla_subset()`: Implementations must return a logical vector of TRUE/FALSE for the subset such that, given the full A matrix and values output, `A[, subset, drop = FALSE]` and `values[subset]` (or `values[subset, , drop = FALSE]` for data.frame values) are equal to the `inla_f = TRUE` version of A and values. The default method uses the `ibm_values` output to construct the subset indexing.
- `ibm_invalid_output()`: Implementations should return a logical vector of length `ibm_n_output(mapper, input, state, ...)` indicating which, if any, output elements of `ibm_eval(mapper, input, state, ...)` are known to be invalid. For for multi/collect mappers, a list, when given a `multi=TRUE` argument.
- `ibm_n(default)`: Returns a non-null element 'n' from the mapper object, and gives an error if it doesn't exist. If `inla_f=TRUE`, first checks for a 'n\_inla' element.
- `ibm_n_output(default)`: Returns `NROW(input)`
- `ibm_values(default)`: Returns a non-null element 'values' from the mapper object, and `seq_len(ibm_n(mapper))` if it doesn't exist.

- `ibm_is_linear(default)`: Returns logical `is_linear` from the mapper object if it exists, and otherwise TRUE.
- `ibm_jacobian(default)`: Mapper classes must implement their own `ibm_jacobian` method.
- `ibm_linear(default)`: Calls `ibm_eval()` and `ibm_jacobian()` and returns a `bm_taylor` object. The `state0` information in the affine mapper indicates for which state the offset was evaluated; The affine mapper output is defined as

```
effect(input, state) =
  offset(input, state0) + jacobian(input, state0) %*% (state - state0)
```

- `ibm_simplify(default)`: Calls `ibm_linear()` for linear mappers, and returns the original mapper for non-linear mappers.
- `ibm_eval(default)`: Verifies that the mapper is linear with `ibm_is_linear()`, and then computes a linear mapping as `ibm_jacobian(...)` %\*% `state`. When `state` is NULL, a zero vector of length `ibm_n_output(...)` is returned.
- `ibm_eval2(default)`: Calls `jacobian <- ibm_jacobian(...)` and `offset <- ibm_eval(..., jacobian = jacobian)` and returns a list with elements `offset` and `jacobian`, as needed by `ibm_linear.default()` and similar methods. Mapper classes can implement their own `ibm_eval2` method if joint construction of evaluation and Jacobian is more efficient than separate or sequential construction.
- `ibm_names(default)`: Returns NULL
- `ibm_inla_subset(default)`: Uses the `ibm_values` output to construct the inla subset indexing, passing extra arguments such as `multi` on to the methods (this means it supports both regular vector values and `multi=1` data.frame values).
- `ibm_invalid_output(default)`: Returns an all-FALSE logical vector.

## See Also

`bru_mapper` for constructor methods, and `bru_get_mapper` for hooks to extract mappers from latent model object class objects.

`bru_mapper`, `bru_get_mapper()`

Other mappers: `bm_aggregate()`, `bm_collect()`, `bm_const()`, `bm_factor()`, `bm_fmesh()`, `bm_harmonics()`, `bm_index()`, `bm_linear()`, `bm_logsumexp()`, `bm_marginal()`, `bm_matrix()`, `bm_mesh_B()`, `bm_multi()`, `bm_pipe()`, `bm_repeat()`, `bm_scale()`, `bm_shift()`, `bm_sum()`, `bm_taylor()`, `bru_get_mapper()`, `bru_mapper()`, `bru_mapper.fm_mesh_1d()`, `bru_mapper.fm_mesh_2d()`

## Examples

```
# ibm_names
mapper <- bm_multi(list(A = bm_index(2), B = bm_index(2)))
ibm_names(mapper)
ibm_names(mapper) <- c("new", "names")
ibm_names(mapper)
```

---

**bru\_model\_mapper\_methods***Mapper methods for model objects*

---

**Description**

Methods for the `ibm_linear()` and `ibm_simplify()` methods for `bru` model objects and related classes.

**Usage**

```
## S3 method for class 'bru_model'
ibm_linear(mapper, input, state = NULL, ...)

## S3 method for class 'bru_comp_list'
ibm_linear(mapper, input, state = NULL, ...)

## S3 method for class 'bru_comp'
ibm_linear(mapper, input, state = NULL, ...)

## S3 method for class 'bru_model'
ibm_simplify(mapper, input = NULL, state = NULL, ...)

## S3 method for class 'bru_comp'
ibm_simplify(mapper, input = NULL, state = NULL, ...)

## S3 method for class 'bru_comp_list'
ibm_simplify(mapper, input = NULL, state = NULL, ...)
```

**Arguments**

<code>mapper</code>	A mapper S3 object, inheriting from <code>bru_mapper</code> .
<code>input</code>	Data input for the mapper.
<code>state</code>	A vector of latent state values for the mapping, of length <code>ibm_n(mapper, inla_f = FALSE)</code>
<code>...</code>	Arguments passed on to other methods

**Functions**

- `ibm_linear(bru_model)`: Returns a list (one element per observation model) of `bm_list` objects, each with one `bm_taylor` entry for each included component.
- `ibm_simplify(bru_model)`: Returns a list (one element per observation model) of `bm_list` objects, each with one `bru_mapper` entry for each included component.

---

**bru\_obs***Observation model construction for usage with bru()*

---

## Description

Observation model construction for usage with [bru\(\)](#).

Note: Prior to version 2.12.0, this function was called `like()`, and that alias will remain for a while until examples etc have been updated and users made aware of the change. The name change is to avoid issues with namespace clashes, e.g. with `data.table::like()`, and also to signal that the function defines observation models, not just likelihood functions.

## Usage

```
bru_obs(  
  formula = . ~ .,  
  family = "gaussian",  
  data = NULL,  
  response_data = NULL,  
  data_extra = NULL,  
  E = NULL,  
  Ntrials = NULL,  
  weights = NULL,  
  scale = NULL,  
  domain = NULL,  
  samplers = NULL,  
  ips = NULL,  
  used = NULL,  
  allow_combine = NULL,  
  aggregate = NULL,  
  aggregate_input = NULL,  
  control.family = NULL,  
  tag = NULL,  
  options = list(),  
  .envir = parent.frame(),  
  include = deprecated(),  
  exclude = deprecated(),  
  include_latent = deprecated()  
)  
  
like(  
  formula = . ~ .,  
  family = "gaussian",  
  data = NULL,  
  response_data = NULL,  
  E = NULL,  
  Ntrials = NULL,
```

```

weights = NULL,
scale = NULL,
domain = NULL,
samplers = NULL,
ips = NULL,
used = NULL,
allow_combine = NULL,
control.family = NULL,
tag = NULL,
options = list(),
.envir = parent.frame(),
mesh = deprecated(),
include = deprecated(),
exclude = deprecated(),
include_latent = deprecated()
)
bru_obs_list(...)

## S3 method for class 'list'
bru_obs_list(object, ..., .envir = NULL)

## S3 method for class 'bru_obs_list'
bru_obs_list(..., .envir = NULL)

## S3 method for class 'bru_obs'
c(..., .envir = NULL)

## S3 method for class 'bru_obs_list'
c(..., .envir = NULL)

## S3 method for class 'bru_obs_list'
x[i]

like_list(...)

bru_like_list(...)

```

## Arguments

- formula** a formula where the right hand side is a general R expression defines the predictor used in the model.
- family** A string identifying a valid INLA::inla likelihood family. The default is gaussian with identity link. In addition to the likelihoods provided by inla (see names(INLA::inla.models())\$lik) inlabru supports fitting latent Gaussian Cox processes via family = "cp". As an alternative to [bru\(\)](#), the [lgcp\(\)](#) function provides a convenient interface to fitting Cox processes.
- data** Predictor expression-specific data, as a `data.frame`, `tibble`, or `sf`. Since

	2.12.0.9023, deprecated support for <code>SpatialPoints[DataFrame]</code> objects.
<code>response_data</code>	Observation/response-specific data for models that need different size/format for inputs and response variables, as a <code>data.frame</code> , <code>tibble</code> , or <code>sf</code> . Since 2.12.0.9023, deprecated support for <code>SpatialPoints[DataFrame]</code> objects.
<code>data_extra</code>	object convertible with <code>as.list()</code> with additional variables to be made available in predictor evaluations. Variables with the same names as the data object will be ignored, unless accessed via <code>.data_extra.[["name"]]</code> or <code>.data_extra.\$name</code> in the formula.
<code>E</code>	Exposure/effort parameter for family = 'poisson' passed on to <code>INLA::inla</code> . Special case if family is 'cp': rescale all integration weights by a scalar E. For sampler specific reweighting/effort, use a weight column in the samplers object instead, see <a href="#">fmesher::fm_int()</a> . Default taken from <code>options\$E</code> , normally 1.
<code>Ntrials</code>	A vector containing the number of trials for the 'binomial' likelihood. Default taken from <code>options\$Ntrials</code> , normally 1.
<code>weights</code>	Fixed (optional) weights parameters of the likelihood, so the <code>log_likelihood[i]</code> is changed into <code>weights[i] * log_likelihood[i]</code> . Default value is 1. WARNING: The normalizing constant for the likelihood is NOT recomputed, so ALL marginals (and the marginal likelihood) must be interpreted with great care. For <code>family = "cp"</code> , the weights are applied as <code>sum(weights * eta)</code> in the point location contribution part of the log-likelihood, where <code>eta</code> is the linear predictor, and do not affect the integration part of the likelihood. This can be used to implement approximative methods for point location uncertainty.
<code>scale</code>	Fixed (optional) scale parameters of the precision for several models, such as Gaussian and student-t response models.
<code>domain, samplers, ips</code>	Arguments used for <code>family="cp"</code> and <code>aggregate=</code> . <code>domain</code> Named list of domain definitions, see <a href="#">fmesher::fm_int()</a> . <code>samplers</code> Integration domain for <code>family="cp"</code> or subdomains for <code>aggregate=</code> , see <a href="#">fmesher::fm_int()</a> . <code>ips</code> Integration points. Defaults to <code>fmesher::fm_int(domain, samplers)</code> . If explicitly given, overrides <code>domain</code> and <code>samplers</code> .
<code>used</code>	Either <code>NULL</code> (default) or a <a href="#">bru_used()</a> object. When, <code>NULL</code> , the information about what effects and latent vectors are made available to the predictor evaluation is defined by <code>bru_used(formula)</code> , which will include all effects and latent vectors used by the predictor expression.
<code>allow_combine</code>	logical; If <code>TRUE</code> , the predictor expression may involve several rows of the input data to influence the same row. When <code>NULL</code> , defaults to <code>FALSE</code> , unless <code>response_data</code> is non- <code>NULL</code> , or <code>data</code> is a <code>list</code> , or the likelihood construction requires it.
<code>aggregate</code>	character ("none", "sum", "average", "logsumexp", or "logaverageexp", as defined by <code>bm_aggregate(type = aggregate)</code> ) or an aggregation <code>bru_mapper</code> object ( <a href="#">bm_aggregate()</a> or <a href="#">bm_logsumexp()</a> ). Default <code>NULL</code> , interpreted as "none". <b>[Experimental]</b> , available from version 2.12.0.9013.

`aggregate_input`  
NULL or an optional input list to the mapper defined by non-NULL aggregate, overriding the default,

```
list(block = .data.[[".block"]],  
     weights = .data.[["weight"]],  
     n_block = bru_response_size(.response_data.))
```

**[Experimental]**, available from version 2.12.0.9013.

`control.family` A optional list of INLA::control.family options

`tag` character; Name that can be used to identify the relevant parts of INLA predictor vector output, via [bru\\_index\(\)](#).

`options` A [bru\\_options](#) options object or a list of options passed on to [bru\\_options\(\)](#)

`.envir` The evaluation environment to use for special arguments (E, Ntrials, weights, and scale) if not found in response\_data or data. Defaults to the calling environment.

`include, exclude, include_latent`  
**[Deprecated]**, use used instead.

`mesh` **[Deprecated]** Ignored.

`...` For `bru_obs_list.bru_obs`, one or more `bru_obs` objects

`object` A list of `bru_obs` objects

`x` `bru_obs_list` object from which to extract element(s)

`i` indices specifying elements to extract

## Value

A likelihood configuration which can be used to parameterise [bru\(\)](#).

## Methods (by generic)

- `c(bru_obs)`: Combine several `bru_obs` objects into a `bru_obs_list` object

## Functions

- `like()`: **[Deprecated]** Legacy `like()` method for `inlabru` prior to version 2.12.0. Use [bru\\_obs\(\)](#) instead.
- `bru_obs_list()`: Combine `bru_obs` observation model object into a `bru_obs_list` object
- `bru_obs_list(list)`: Combine one or more lists of `bru_obs` observation model objects into a `bru_obs_list` object
- `bru_obs_list(bru_obs_list)`: Combine a list of `bru_obs` observation model objects into a `bru_obs_list` object
- `c(bru_obs_list)`: Combine several `bru_obs_list` objects into a `bru_obs_list` object
- `like_list()`: **[Deprecated]** Backwards compatibility for versions <= 2.12.0. For later versions, use `as_bru_obs_list()`, `bru_obs_list()`, or `c()`.
- `bru_like_list()`: **[Deprecated]** Backwards compatibility for versions <= 2.12.0.9017. For later versions, use `as_bru_obs_list()`, `bru_obs_list()` or `c()`.

**Author(s)**

Fabian E. Bachl <bachlfab@gmail.com>  
 Finn Lindgren <finn.lindgren@gmail.com>

**See Also**

[bru\\_response\\_size\(\)](#), [bru\\_used\(\)](#), [bru\\_comp\(\)](#), [bru\\_comp\\_eval\(\)](#)  
[summary.bru\\_obs\(\)](#)

**Examples**

```
if (bru_safe_inla() &&
    require(ggplot2, quietly = TRUE)) {

  # The 'bru_obs()' (previously 'like()') function's main purpose is to set up
  # observation models, both for single- and multi-likelihood models.
  # The following example generates some random covariates which are observed
  # through two different random effect models with different likelihoods

  # Generate the data

  set.seed(123)

  n1 <- 200
  n2 <- 10

  x1 <- runif(n1)
  x2 <- runif(n2)
  z2 <- runif(n2)

  y1 <- rnorm(n1, mean = 2 * x1 + 3)
  y2 <- rpois(n2, lambda = exp(2 * x2 + z2 + 3))

  df1 <- data.frame(y = y1, x = x1)
  df2 <- data.frame(y = y2, x = x2, z = z2)

  # Single likelihood models and inference using bru are done via

  cmp1 <- y ~ -1 + Intercept(1) + x
  fit1 <- bru(cmp1, family = "gaussian", data = df1)
  summary(fit1)

  cmp2 <- y ~ -1 + Intercept(1) + x + z
  fit2 <- bru(cmp2, family = "poisson", data = df2)
  summary(fit2)

  # A joint model has two likelihoods, which are set up using the bru_obs
  # function

  lik1 <- bru_obs(
    "gaussian",
```

```

    formula = y ~ x + Intercept,
    data = df1,
    tag = "norm"
  )
lik2 <- bru_obs(
  "poisson",
  formula = y ~ x + z + Intercept,
  data = df2,
  tag = "pois"
)
# The union of effects of both models gives the components needed to run bru

jcmp <- ~ x + z + Intercept(1)
jfit <- bru(jcmp, lik1, lik2)

bru_index(jfit, "norm")
bru_index(jfit, "pois")

# Compare the estimates

p1 <- ggplot() +
  gg(fit1$summary.fixed, bar = TRUE) +
  ylim(0, 4) +
  ggtitle("Model 1")
p2 <- ggplot() +
  gg(fit2$summary.fixed, bar = TRUE) +
  ylim(0, 4) +
  ggtitle("Model 2")
pj <- ggplot() +
  gg(jfit$summary.fixed, bar = TRUE) +
  ylim(0, 4) +
  ggtitle("Joint model")

multiplot(p1, p2, pj)
}

```

**bru\_options***Create or update an options objects***Description**

Create a new options object, or merge information from several objects.

The `_get`, `_set`, and `_reset` functions operate on a global package options override object. In many cases, setting options in specific calls to `bru()` is recommended instead.

**Usage**

```
bru_options(...)

as.bru_options(x = NULL)

bru_options_default()

bru_options_check(options, ignore_null = TRUE)

bru_options_get(name = NULL, include_default = TRUE)

bru_options_set(..., .reset = FALSE)

bru_options_reset()

bru_options_set_local(..., .reset = FALSE, .envir = parent.frame())
```

**Arguments**

...	A collection of named options, optionally including one or more <code>bru_options</code> objects. Options specified later override the previous options.
x	An object to be converted to an <code>bru_options</code> object.
options	An <code>bru_options</code> object to be checked
ignore_null	Ignore missing or NULL options.
name	Either NULL, or single option name string, or character vector or list with option names, Default: NULL
include_default	logical; If TRUE, the default options are included together with the global override options. Default: TRUE
.reset	For <code>bru_options_set</code> , logical indicating if the global override options list should be emptied before setting the new option(s).
.envir	The environment in which to set the options. Default: <code>parent.frame()</code>

**Value**

`bru_options()` returns a `bru_options` object.

For `as.bru_options()`, NULL or no input returns an empty `bru_options` object, a list is converted via `bru_options(...)`, and `bru_options` input is passed through. Other types of input generates an error.

`bru_options_default()` returns an `bru_options` object containing default options.

`bru_options_check()` returns a logical; TRUE if the object contains valid options for use by other functions

`bru_options_get` returns either an `bru_options` object, for `name == NULL`, the contents of single option, if `name` is a options name string, or a named list of option contents, if `name` is a list of option name strings.

`bru_options_set()` returns a copy of the global override options, invisibly (as `bru_options_get(include_default = FALSE)`).

## Functions

- `as.bru_options()`: Coerces inputs to a `bru_options` object.
- `bru_options_default()`: Returns the default options.
- `bru_options_check()`: Checks for valid contents of a `bru_options` object, and produces warnings for invalid options.
- `bru_options_get()`: Used to access global package options.
- `bru_options_set()`: Used to set global package options.
- `bru_options_reset()`: Clears the global option overrides.
- `bru_options_set_local()`: Sets local option overrides, that are automatically reset using `withr::defer()`.

## Valid options

For `bru_options` and `bru_options_set`, recognised options are:

**bru\_verbose** logical or numeric; if TRUE, log messages of verbosity  $\leq 1$  are printed by [bru\\_log\\_message\(\)](#).

If numeric, log messages of verbosity  $\leq \text{bru_verbose}$  are printed. For line search details, set `bru_verbose=2` or 3. Default: 0, to not print any messages

**bru\_verbose\_store** logical or numeric; if TRUE, log messages of verbosity  $\leq 1$  are stored by [bru\\_log\\_message\(\)](#). If numeric, log messages of verbosity  $\leq$  are stored. Default: Inf, to store all messages.

**bru\_run** If TRUE, run inference. Otherwise only return configuration needed to run inference.

**bru\_max\_iter** maximum number of inla iterations, default 10. Also see the `bru_method$rel_tol` and related options below.

**bru\_initial** An `inla` object returned from previous calls of `INLA::inla`, [bru\(\)](#) or [lgcp\(\)](#), or a list of named vectors of starting values for the latent variables. This will be used as a starting point for further improvement of the approximate posterior.

**bru\_int\_args** List of arguments passed all the way to the integration method `ipoints` and `int.polygon` for 'cp' family models;

**method** "stable" or "direct". For "stable" (default) integration points are aggregated to mesh vertices.

**nsub1** Number of integration points per knot interval in 1D. Default 30.

**nsub2** Number of integration points along a triangle edge for 2D. Default 9.

**nsub** Deprecated parameter that overrides `nsub1` and `nsub2` if set. Default NULL.

**bru\_method** List of arguments controlling the iterative inlabru method:

**taylor** 'pandemic' (default, from version 2.1.15).

**search** Either 'all' (default), to use all available line search methods, or one or more of

'finite' (reduce step size until predictor is finite)

'contract' (decrease step size until trust hypersphere reached)

'expand' (increase step size until no improvement)

**'optimise'** (fast approximate error norm minimisation)

To disable line search, set to an empty vector. Line search is not available for taylor="legacy".

**factor** Numeric,  $> 1$  determining the line search step scaling multiplier. Default  $(1 + \sqrt{5})/2$ .

**rel\_tol** Stop the iterations when the largest change in linearisation point (the conditional latent state mode) in relation to the estimated posterior standard deviation is less than **rel\_tol**. Default 0.1 (ten percent).

**max\_step** The largest allowed line search step factor. Factor 1 is the full INLA step. Default is 2.

**line\_opt\_method** Which method to use for the line search optimisation step. Default "onestep", using a quadratic approximation based on the value and gradient at zero, and the value at the current best step length guess. The method "full" does line optimisation on the full nonlinear predictor; this is slow and intended for debugging purposes only.

**bru\_compress\_cp** logical; when TRUE, compress the  $\sum_{i=1}^n \eta_i$  part of the Poisson process likelihood (family="cp") into a single term, with  $y = n$ , and predictor mean(eta). Default: TRUE

**bru\_debug** logical; when TRUE, activate temporary debug features for package development. Default: FALSE

**inla() options** All options not starting with bru\_ are passed on to inla(), sometimes after altering according to the needs of the inlabru method. Warning: Due to how inlabru currently constructs the inla() call, the mean, prec, mean.intercept, and prec.intercept settings in control.fixed will have no effect. Until a more elegant alternative has been implemented, use explicit mean.linear and prec.linear specifications in each model="linear" component instead.

## See Also

[bru\\_options\(\)](#), [bru\\_options\\_default\(\)](#), [bru\\_options\\_get\(\)](#)

## Examples

```
## Not run:
if (interactive()) {
  # Combine global and user options:
  options1 <- bru_options(bru_options_get(), bru_verbose = TRUE)
  # Create a proto-options object in two equivalent ways:
  options2 <- as.bru_options(bru_verbose = TRUE)
  options2 <- as.bru_options(list(bru_verbose = TRUE))
  # Combine options objects:
  options3 <- bru_options(options1, options2)
}

## End(Not run)
## Not run:
if (interactive()) {
  bru_options_check(bru_options(bru_max_iter = "text"))
}

## End(Not run)
bru_options_get("bru_verbose")
```

```

## Not run:
if (interactive()) {
  bru_options_set(
    bru_verbose = TRUE,
    verbose = TRUE
  )
}

## End(Not run)
my_fun <- function(val) {
  bru_options_set_local(bru_verbose = val)
  bru_options_get("bru_verbose")
}
# Inside the function, the bru_verbose option is changed.
# Outside the function, the bru_verbose option is unchanged.
print(my_fun(TRUE))
print(bru_options_get("bru_verbose"))
print(my_fun(FALSE))
print(bru_options_get("bru_verbose"))

```

**bru\_response\_size**      *Response size queries*

### Description

Extract the number of response values from `bru` and related objects.

### Usage

```

bru_response_size(object)

## Default S3 method:
bru_response_size(object)

## S3 method for class 'inla.surv'
bru_response_size(object)

## S3 method for class 'bru_obs'
bru_response_size(object)

## S3 method for class 'bru_obs_list'
bru_response_size(object)

## S3 method for class 'bru_info'
bru_response_size(object)

## S3 method for class 'bru'
bru_response_size(object)

```

**Arguments**

object	An object from which to extract response size(s).
--------	---

**Value**

An integer vector.

**Methods (by class)**

- `bru_response_size(default)`: Extract the number of observations from an object supporting `NROW()`.
- `bru_response_size(inla.surv)`: Extract the number of observations from an `inla.surv` object.
- `bru_response_size(bru_obs)`: Extract the number of observations from a `bru_obs` object.
- `bru_response_size(bru_obs_list)`: Extract the number of observations from a `bru_obs_list` object, as a vector with one value per observation model.
- `bru_response_size(bru_info)`: Extract the number of observations from a `bru_info` object, as a vector with one value per observation model.
- `bru_response_size(bru)`: Extract the number of observations from a `bru` object, as a vector with one value per observation model.

**See Also**

[like\(\)](#)

**Examples**

```
bru_response_size(  
  bru_obs(y ~ 1, data = data.frame(y = rnorm(10)), family = "gaussian")  
)
```

---

`bru_set_missing`      *Set missing values in observation models*

---

**Description**

Set all or parts of the observation model response data to NA, for example for use in cross validation (with [bru\\_rerun\(\)](#)) or prior sampling (with [bru\\_rerun\(\)](#) and [generate\(\)](#)).

## Usage

```
bru_set_missing(object, keep = FALSE, ...)
## S3 method for class 'bru'
bru_set_missing(object, keep = FALSE, ...)

## S3 method for class 'bru_obs_list'
bru_set_missing(object, keep = FALSE, ...)

## S3 method for class 'bru_obs'
bru_set_missing(object, keep = FALSE, ...)
```

## Arguments

object	A <i>bru</i> , <i>bru_obs</i> or <i>bru_obs_list</i> object
keep	For <i>bru_obs</i> , a single logical or an integer vector; If TRUE, keep all the response data, if FALSE (default), set all of it to NA. An integer vector determines which elements to keep (for positive values) or to set as missing (negative values). For <i>bru</i> and <i>bru_obs_list</i> , a logical scalar or vector, or a list, see Details.
...	Additional arguments passed on to the <i>bru_obs</i> method. Currently unused.

## Details

For *bru* and *bru\_obs\_list*,

- *keep* must be either a single logical, which is expanded to a list,
- a logical vector, which is converted to a list,
- an unnamed list of the same length as the number of observation models, with elements compatible with the *bru\_obs* method, or
- a named list with elements compatible with the *bru\_obs* method, and only the named *bro\_obs* models are acted upon, i.e. the elements not present in the list are treated as *keep* = TRUE.

E.g.: *keep* = *list*(*b* = FALSE) sets all observations in model *b* to missing, and does not change model *a*.

E.g.: *keep* = *list*(*a* = 1:4, *b* = -(3:5)) keeps only observations 1:4 of model *a*, marking the rest as missing, and sets observations 3:5 of model *b* to missing.

## Examples

```
obs <- c(
  A = bru_obs(y_A ~ ., data = data.frame(y_A = 1:6)),
  B = bru_obs(y_B ~ ., data = data.frame(y_B = 11:15))
)
bru_response_size(obs)
lapply(
  bru_set_missing(obs, keep = FALSE),
  function(x) {
    x[["response_data"]][["BRU_response"]]
```

```

        }
    )
lapply(
  bru_set_missing(obs, keep = list(B = FALSE)),
  function(x) {
    x[["response_data"]][["BRU_response"]]
  }
)
lapply(
  bru_set_missing(obs, keep = list(1:4, -(3:5))),
  function(x) {
    x[["response_data"]][["BRU_response"]]
  }
)

```

**bru\_timings***Extract timing information from fitted [bru](#) object***Description**

Extracts a data.frame or tibble with information about the Time (CPU), System, and Elapsed time for each step of a `bru()` run.

**Usage**

```

bru_timings(object, ...)
## S3 method for class 'bru'
bru_timings(object, ...)

```

**Arguments**

<code>object</code>	A fitted <code>bru</code> object
<code>...</code>	unused

**bru\_timings\_plot***Plot inlabru iteration timings***Description**

Draws the time per iteration for preprocessing (including linearisation), `inla()` calls, and line search. Iteration 0 is the time used for defining the model structure.

**Usage**

```
bru_timings_plot(x)
```

## Arguments

- x a **bru** object, typically a result from **bru()** for a nonlinear predictor model

## Details

Requires the "ggplot2" package to be installed.

## Examples

```
## Not run:
fit <- bru(...)
bru_timings_plot(fit)

## End(Not run)
```

**bru\_transformation**      *Transformation tools*

## Description

Tools for transforming between N(0,1) variables and other distributions in predictor expressions

## Usage

```
bru_forward_transformation(qfun, x, ..., tail.split. = 0)
bru_inverse_transformation(pfun, x, ..., tail.split. = NULL)
```

## Arguments

- |             |   |
|-------------|---|
| qfun        | A quantile function object, such as qexp  |
| x           | Values to be transformed  |
| ...         | Distribution parameters passed on to the qfun and pfun functions  |
| tail.split. | For x-values larger than tail.split., upper quantile calculations are used internally, and for smaller values lower quantile calculations are used. This can avoid lack of accuracy in the distribution tails. If NULL, forward calculations split at 0, and inverse calculations use lower tails only, potentially losing accuracy in the upper tails. |
| pfun        | A CDF function object, such as pexp   |

## Value

- For **bru\_forward\_transformation**, a numeric vector
- For **bru\_inverse\_transformation**, a numeric vector

## Examples

```
u <- rnorm(5, 0, 1)
y <- bru_forward_transformation(qexp, u, rate = 2)
v <- bru_inverse_transformation(pexp, y, rate = 2)
rbind(u, y, v)
```

deltaIC

*Summarise DIC and WAIC from lgcp objects.*

## Description

Calculates DIC and/or WAIC differences and produces an ordered summary.

## Usage

```
deltaIC(..., criterion = "DIC")
```

## Arguments

- |           |  |
|-----------|--|
| ...       | Comma-separated objects inheriting from class <code>inla</code> and obtained from a run of <code>INLA::inla()</code> , <code>bru()</code> or <code>lgcp()</code> |
| criterion | character vector. If it includes 'DIC', computes DIC differences; If it contains 'WAIC', computes WAIC differences. Default: 'DIC'                               |

## Value

A data frame with each row containing the Model name, DIC and Delta.DIC, and/or WAIC and Delta.WAIC.

## Examples

```
if (bru_safe_inla()) {
  # Generate some data
  input.df <- data.frame(idx = 1:10, x = cos(1:10))
  input.df <- within(
    input.df,
    y <- rpois(10, 5 + 2 * cos(1:10) + rnorm(10, mean = 0, sd = 0.1))
  )

  # Fit two models
  fit1 <- bru(
    y ~ x,
    family = "poisson",
    data = input.df,
    options = list(control.compute = list(dic = TRUE))
  )
  fit2 <- bru(
    y ~ x + rand(idx, model = "iid"),
```

```

family = "poisson",
data = input.df,
options = list(control.compute = list(dic = TRUE))
)

# Compare DIC

deltaIC(fit1, fit2)
}

```

**devel.cvmeasure***Variance and correlations measures for prediction components***Description**

Calculates local and integrated variance and correlation measures as introduced by Yuan et al. (2017).

**Usage**

```
devel.cvmeasure(joint, prediction1, prediction2, samplers = NULL, mesh = NULL)
```

**Arguments**

- `joint` A joint prediction of two latent model components.
- `prediction1` A prediction of the first component.
- `prediction2` A prediction of the second component.
- `samplers` An sf or SpatialPolygon object describing the area for which to compute the cumulative variance measure.
- `mesh` The `fmesher::fm_mesh_2d` for which the prediction was performed (required for cumulative Vmeasure).

**Value**

Variance and correlations measures.

**References**

Y. Yuan, F. E. Bachl, F. Lindgren, D. L. Brochers, J. B. Illian, S. T. Buckland, H. Rue, T. Gerrodette. 2017. Point process models for spatio-temporal distance sampling data from a large-scale survey of blue whales. <https://arxiv.org/abs/1604.06013>

## Examples

```

if (bru_safe_inla() &&
    require("ggplot2", quietly = TRUE) &&
    require("patchwork", quietly = TRUE) &&
    require("sn", quietly = TRUE) &&
    require("terra", quietly = TRUE) &&
    require("sf", quietly = TRUE) &&
    require("RColorBrewer", quietly = TRUE) &&
    require("dplyr", quietly = TRUE) &&
    require("magrittr", quietly = TRUE)) {
  # Load Gorilla data

  gorillas <- gorillas_sf
  gorillas$gcov <- gorillas_sf_gcov()

  # Use RColorBrewer

  # Fit a model with two components:
  # 1) A spatial smooth SPDE
  # 2) A spatial covariate effect (vegetation)

  pcmatern <- INLA::inla.spde2.pcmatern(
    gorillas$mesh,
    prior.sigma = c(0.1, 0.01),
    prior.range = c(0.01, 0.01)
  )

  cmp <- geometry ~ 0 +
    vegetation(gorillas$gcov$vegetation, model = "factor_contrast") +
    spde(geometry, model = pcmatern) -
    Intercept(1)

  fit <- lgcp(
    cmp,
    gorillas$nests,
    samplers = gorillas$boundary,
    domain = list(geometry = gorillas$mesh),
    options = list(control.inla = list(int.strategy = "eb")))
}

# Predict SPDE and vegetation at a grid covering the domain of interest
pred_loc <- fm_pixels(
  gorillas$mesh,
  mask = gorillas$boundary,
  dims = c(200, 200),
  format = "sf"
)
pred <- predict(
  fit,
  pred_loc,
  ~ list(
    joint = spde + vegetation,

```

```

        field = spde,
        veg = vegetation
    )
)

pred_collect <-
  rbind(
    pred$joint %>% dplyr::mutate(component = "joint"),
    pred$field %>% dplyr::mutate(component = "field"),
    pred$veg %>% dplyr::mutate(component = "veg")
  ) %>%
  dplyr::mutate(var = sd^2)

# Plot component mean

ggplot(pred_collect) +
  geom_tile(aes(geometry = geometry, fill = mean), stat = "sf_coordinates") +
  coord_equal() +
  facet_wrap(~component, nrow = 1) +
  theme(legend.position = "bottom")

# Plot component variance

ggplot(pred_collect) +
  geom_tile(aes(geometry = geometry, fill = var), stat = "sf_coordinates") +
  coord_equal() +
  facet_wrap(~component, nrow = 1) +
  theme(legend.position = "bottom")

# Calculate variance and correlation measure

vm <- devel.cvmeasure(pred$joint, pred$field, pred$veg)
# Compute nominal relative variance contributions; note that these can be
# greater than 100%
vm <- dplyr::mutate(
  vm,
  var1_rel = if_else(var1 <= 0, NA, var1 / var.joint),
  var2_rel = if_else(var2 <= 0, NA, var2 / var.joint)
)
lprange <- range(vm$var.joint, vm$var1, vm$var2)
vm <- tidyverse::pivot_longer(vm,
  cols = c(var.joint, var1, var2, cov, cor, var1_rel, var2_rel),
  names_to = "component",
  values_to = "value"
)

# Variance contribution of the components

csc <- scale_fill_gradientn(
  colours = brewer.pal(9, "YlOrRd"),
  limits = lprange
)

```

```

vm_ <- dplyr::filter(vm, component %in% c("var.joint", "var1", "var2"))
ggplot(vm_) +
  geom_tile(aes(geometry = geometry, fill = value),
            stat = "sf_coordinates"
  ) +
  csc +
  coord_equal() +
  facet_wrap(~component, nrow = 1) +
  theme(legend.position = "bottom")

# Relative variance contribution of the components
# When bo

vm_ <- dplyr::filter(vm, component %in% c("var1_rel", "var2_rel"))
ggplot(vm_) +
  geom_tile(aes(geometry = geometry, fill = value),
            stat = "sf_coordinates"
  ) +
  scale_fill_gradientn(
    colours = brewer.pal(9, "YlOrRd"),
    trans = "log"
  ) +
  coord_equal() +
  facet_wrap(~component, nrow = 1)

# Where both relative contributions are larger than 1, the posterior
# correlations are strongly negative.

# Covariance and correlation of field and vegetation

vm_cov <- dplyr::filter(vm, component %in% "cov")
vm_cor <- dplyr::filter(vm, component %in% "cor")
(ggplot(vm_cov) +
  geom_tile(aes(geometry = geometry, fill = value),
            stat = "sf_coordinates"
  ) +
  scale_fill_gradientn(
    colours = rev(brewer.pal(9, "RdBu")),
    limits = c(-1, 1) * max(abs(vm_cov$value), na.rm = TRUE)
  ) +
  coord_equal() +
  theme(legend.position = "bottom") +
  ggtitle("Covariances") |
  ggplot(vm_cor) +
  geom_tile(aes(geometry = geometry, fill = value),
            stat = "sf_coordinates"
  ) +
  scale_fill_gradientn(
    colours = rev(brewer.pal(9, "RdBu")),
    limits = c(-1, 1) * max(abs(vm_cor$value), na.rm = TRUE)
  ) +
  coord_equal() +
  theme(legend.position = "bottom") +

```

```

    ggtile("Correlations")
  )

# Variance and correlation integrated over space

vrt <- fm_vertices(gorillas$mesh, format = "sf")
pred_vrt <- predict(
  fit,
  vrt,
  ~ list(
    joint = spde + vegetation,
    field = spde,
    veg = vegetation
  )
)

vm.int <- devel.cvmeasure(
  pred_vrt$joint,
  pred_vrt$field,
  pred_vrt$veg,
  samplers = fm_int(gorillas$mesh, gorillas$boundary),
  mesh = gorillas$mesh
)
vm.int
}

```

**eval\_spatial**      *Evaluate spatial covariates*

## Description

Evaluate spatial covariates

## Usage

```

eval_spatial(data, where, layer = NULL, selector = NULL)

## S3 method for class 'SpatialPolygonsDataFrame'
eval_spatial(data, where, layer = NULL, selector = NULL)

## S3 method for class 'SpatialPixelsDataFrame'
eval_spatial(data, where, layer = NULL, selector = NULL)

## S3 method for class 'SpatialGridDataFrame'
eval_spatial(data, where, layer = NULL, selector = NULL)

## S3 method for class 'sf'
eval_spatial(data, where, layer = NULL, selector = NULL)

```

```
## S3 method for class 'SpatRaster'
eval_spatial(data, where, layer = NULL, selector = NULL)

## S3 method for class 'stars'
eval_spatial(data, where, layer = NULL, selector = NULL)
```

**Arguments**

data	Spatial data
where	Where to evaluate the data
layer	Which data layer to extract (as integer or character). May be a vector, specifying a separate layer for each where item.
selector	The name of a variable in where specifying the layer information.

**Methods (by class)**

- eval\_spatial(SpatialPolygonsDataFrame): Compatibility wrapper for eval\_spatial.sf
- eval\_spatial(sf): Supports point-in-polygon information lookup. Other combinations are untested.

format.bru\_mapper      *mapper object summaries*

**Description**

mapper object summaries

**Usage**

```
## S3 method for class 'bru_mapper'
format(x, ..., prefix = "", initial = prefix, depth = 1)

## S3 method for class 'bm_list'
format(
  x,
  ...,
  prefix = "",
  initial = prefix,
  depth = 1,
  collapse = ", ",
  labels = TRUE
)

## S3 method for class 'bru_mapper'
summary(object, ..., prefix = "", initial = prefix, depth = 1)
```

```

## S3 method for class 'bm_multi'
format(x, ..., prefix = "", initial = prefix, depth = 1)

## S3 method for class 'bm_pipe'
format(x, ..., prefix = "", initial = prefix, depth = 1)

## S3 method for class 'bm_collect'
format(x, ..., prefix = "", initial = prefix, depth = 1)

## S3 method for class 'bm_sum'
format(x, ..., prefix = "", initial = prefix, depth = 1)

## S3 method for class 'bm_repeat'
format(x, ..., prefix = "", initial = prefix, depth = 1)

## S3 method for class 'summary_bru_mapper'
print(x, ..., sep = "\n")

## S3 method for class 'bru_mapper'
print(x, ..., sep = "\n", prefix = "", initial = prefix, depth = 1)

## S3 method for class 'bm_list'
print(
  x,
  ...,
  sep = "\n",
  prefix = "",
  initial = prefix,
  depth = 1,
  labels = TRUE,
  collapse = ", "
)

```

## Arguments

x	Object to format/print
...	Unused arguments
prefix	character prefix for each line. Default "".
initial	character prefix for the first line. Default initial=prefix.
depth	The recursion depth for multi/collection/pipe mappers. Default 1, to only show the collection, and not the contents of the sub-mappers.
collapse	character or NULL, as in <code>base:::paste()</code> .
labels	logical; if TRUE, include mapper names or numerical indices. Default TRUE
object	Object to summarise
sep	character; separator for printing the summary.

## Examples

```

mapper <-
  bm_pipe(
    list(
      bm_multi(list(
        A = bm_index(2),
        B = bm_index(3)
      )),
      bm_index(2)
    )
  )
summary(mapper, depth = 2)
mapper <-
  bm_repeat(
    bm_multi(
      list(
        A = bm_index(2),
        B = bm_index(3)
      )
    ),
    3
  )
summary(mapper)
summary(mapper, depth = 0)

```

generate

*Generate samples from fitted bru models*

## Description

Generic function for sampling for fitted models. The function invokes particular methods which depend on the class of the first argument.

Takes a fitted bru object produced by the function [bru\(\)](#) and produces samples given a new set of values for the model covariates or the original values used for the model fit. The samples can be based on any R expression that is valid given these values/covariates and the joint posterior of the estimated random effects.

## Usage

```

generate(object, ...)

## S3 method for class 'bru'
generate(
  object,
  newdata = NULL,
  formula = NULL,
  n.samples = 100,
  seed = 0L,

```

```

  num.threads = NULL,
  used = NULL,
  ...,
  data = deprecated(),
  include = deprecated(),
  exclude = deprecated()
)

```

## Arguments

object	A bru object obtained by calling <a href="#">bru()</a> .
...	additional, unused arguments.
newdata	A data.frame or SpatialPointsDataFrame of covariates needed for sampling.
formula	A formula where the right hand side defines an R expression to evaluate for each generated sample. If NULL, the latent and hyperparameter states are returned as named list elements. See Details for more information.
n.samples	Integer setting the number of samples to draw in order to calculate the posterior statistics. The default, 100, is rather low but provides a quick approximate result.
seed	Random number generator seed passed on to INLA::inla.posterior.sample
num.threads	Specification of desired number of threads for parallel computations. Default NULL, leaves it up to INLA. When seed != 0, overridden to "1:1"
used	Either NULL or a <a href="#">bru_used()</a> object. Default, NULL, uses auto-detection of used variables in the formula.
data	<b>[Deprecated]</b> Use newdata instead.
include, exclude	<b>[Deprecated]</b> If auto-detection of used variables fails, use used instead.

## Details

In addition to the component names (that give the effect of each component evaluated for the input data), the suffix \_latent variable name can be used to directly access the latent state for a component, and the suffix function \_eval can be used to evaluate a component at other input values than the expressions defined in the component definition itself, e.g. `field_eval(cbind(x, y))` for a component that was defined with `field(coordinates, ...)` (see also [bru\\_comp\\_eval\(\)](#)).

For "iid" models with `mapper = bm_index(n)`, `rnorm()` is used to generate new realisations for indices greater than n.

## Value

The form of the value returned by `generate()` depends on the data class and prediction formula. Normally, a data.frame is returned, or a list of data.frames (if the prediction formula generates a list)

List of generated samples

**See Also**

[predict.bru](#)

**Examples**

```
if (bru_safe_inla() &&
    require("sn", quietly = TRUE)) {

  # Generate data for a simple linear model

  input.df <- data.frame(x = cos(1:10))
  input.df <- within(input.df, y <- 5 + 2 * cos(1:10) + rnorm(10, mean = 0, sd = 0.1))

  # Fit the model

  fit <- bru(y ~ xeff(main = x, model = "linear"),
             family = "gaussian", data = input.df
  )
  summary(fit)

  # Generate samples for some predefined x

  df <- data.frame(x = seq(-4, 4, by = 0.1))
  smp <- generate(fit, df, ~ xeff + Intercept, n.samples = 10)

  # Plot the resulting realizations

  plot(df$x, smp[, 1], type = "l")
  for (k in 2:ncol(smp)) points(df$x, smp[, k], type = "l")

  # We can also draw samples from the joint posterior

  df <- data.frame(x = 1)
  smp <- generate(fit, df, ~ data.frame(xeff, Intercept), n.samples = 10)
  smp[[1]]

  # ... and plot them
  if (require(ggplot2, quietly = TRUE)) {
    plot(do.call(rbind, smp))
  }
}
```

**Description**

gg is a generic function for generating geometes from various kinds of spatial objects, e.g. Spatial\* data, meshes, Raster objects and inla/inlabru predictions. The function invokes particular methods which depend on the [class](#) of the first argument.

## Usage

```
gg(data, ...)
```

## Arguments

- data           an object for which to generate a geom.
- ...            Arguments passed on to the geom method.

## Value

The form of the value returned by `gg` depends on the class of its argument. See the documentation of the particular methods for details of what is produced by that method.

## See Also

- Other geomes for inla and inlabru predictions: [gg.bru\\_prediction\(\)](#), [gg.data.frame\(\)](#), [gg.matrix\(\)](#)
- Other geomes for spatial data: [gg.SpatRaster\(\)](#), [gg.SpatialGridDataFrame\(\)](#), [gg.SpatialLines\(\)](#), [gg.SpatialPixels\(\)](#), [gg.SpatialPixelsDataFrame\(\)](#), [gg.SpatialPoints\(\)](#), [gg.SpatialPolygons\(\)](#), [gg.sf\(\)](#)
- Other geomes for meshes: [gg.fm\\_mesh\\_1d\(\)](#), [gg.fm\\_mesh\\_2d\(\)](#)
- Other geomes for Raster data: [gg.RasterLayer\(\)](#)

## Examples

```
if (require("ggplot2", quietly = TRUE)) {
  # Load Gorilla data

  gorillas <- inlabru::gorillas_sf

  # Invoke ggplot and add geomes for the Gorilla nests and the survey
  # boundary

  ggplot() +
    gg(gorillas$boundary) +
    gg(gorillas$nests)
}
```

**gg.bru\_prediction**      *Geom for predictions*

## Description

This geom serves to visualize prediction objects which usually results from a call to [predict.bru\(\)](#). Predictions objects provide summary statistics (mean, median, sd, ...) for one or more random variables. For single variables (or if requested so by setting `bar = TRUE`), a boxplot-style geom

is constructed to show the statistics. For multivariate predictions the mean of each variable (y-axis) is plotted against the row number of the variable in the prediction data frame (x-axis) using `geom_line`. In addition, a `geom_ribbon` is used to show the confidence interval.

Note: `gg.bru_prediction` also understands the format of INLA-style posterior summaries, e.g. `fit$summary.fixed` for an `inla` object `fit`

Requires the `ggplot2` package.

## Usage

```
## S3 method for class 'bru_prediction'
gg(data, mapping = NULL, ribbon = TRUE, alpha = NULL, bar = FALSE, ...)

## S3 method for class 'prediction'
gg(data, ...)
```

## Arguments

<code>data</code>	A prediction object, usually the result of a <code>predict.bru()</code> call.
<code>mapping</code>	a set of aesthetic mappings created by <code>aes</code> . These are passed on to <code>geom_line()</code> . If "fill" is present, it is passed to <code>geom_ribbon()</code> .
<code>ribbon</code>	If TRUE, plot a ribbon around the line based on the smallest and largest quantiles present in the data, found by matching names starting with <code>q</code> and followed by a numerical value. <code>inla()</code> -style numeric+"quant" names are converted to <code>inlabru</code> style before matching.
<code>alpha</code>	The ribbons numeric alpha (transparency) level in [0,1].
<code>bar</code>	If TRUE plot boxplot-style summary for each variable.
...	Arguments passed on to <code>geom_line</code> .

## Value

Concatenation of a `geom_line` value and optionally a `geom_ribbon` value.

## See Also

Other geomes for `inla` and `inlabru` predictions: `gg()`, `gg.data.frame()`, `gg.matrix()`

## Examples

```
if (bru_safe_inla() &&
    require(sn, quietly = TRUE) &&
    require(ggplot2, quietly = TRUE)) {
  # Generate some data

  input.df <- data.frame(x = cos(1:10))
  input.df <- within(input.df, y <- 5 + 2 * cos(1:10) + rnorm(10, mean = 0, sd = 0.1))

  # Fit a model with fixed effect 'x' and intercept 'Intercept'
```

```

fit <- bru(y ~ x, family = "gaussian", data = input.df)

# Predict posterior statistics of 'x'

xpost <- predict(fit, NULL, formula = ~x_latent)

# The statistics include mean, standard deviation, the 2.5% quantile, the median,
# the 97.5% quantile, minimum and maximum sample drawn from the posterior as well as
# the coefficient of variation and the variance.

xpost

# For a single variable like 'x' the default plotting method invoked by gg() will
# show these statistics in a fashion similar to a box plot:
ggplot() +
  gg(xpost)

# The predict function can also be used to simultaneously estimate posteriors
# of multiple variables:

xipost <- predict(fit,
  newdata = NULL,
  formula = ~ c(
    Intercept = Intercept_latent,
    x = x_latent
  )
)
xipost

# If we still want a plot in the previous style we have to set the bar parameter to TRUE

p1 <- ggplot() +
  gg(xipost, bar = TRUE)
p1

# Note that gg also understands the posterior estimates generated while running INLA

p2 <- ggplot() +
  gg(fit$summary.fixed, bar = TRUE)
multiplot(p1, p2)

# By default, if the prediction has more than one row, gg will plot the column 'mean' against
# the row index. This is for instance useful for predicting and plotting function
# but not very meaningful given the above example:

ggplot() +
  gg(xipost)

# For ease of use we can also type

plot(xipost)

```

```
# This type of plot will show a ribbon around the mean, which visualizes the upper and lower  
# quantiles mentioned above (2.5 and 97.5%). Plotting the ribbon can be turned off using the  
# \code{ribbon} parameter  
  
ggplot() +  
  gg(xipost, ribbon = FALSE)  
  
# Much like the other geoms produced by gg we can adjust the plot using ggplot2 style  
# commands, for instance  
  
ggplot() +  
  gg(xipost) +  
  gg(xipost, mapping = aes(y = median), ribbon = FALSE, color = "red")  
}
```

---

**gg.data.frame***Geom for data.frame*

---

**Description**

This geom constructor will simply call [gg.bru\\_prediction\(\)](#) for the data provided.

**Usage**

```
## S3 method for class 'data.frame'  
gg(...)
```

**Arguments**

...              Arguments passed on to [gg.bru\\_prediction\(\)](#).

**Details**

Requires the ggplot2 package.

**Value**

Concatenation of a geom\_line value and optionally a geom\_ribbon value.

**See Also**

Other geoms for inla and inlabru predictions: [gg\(\)](#), [gg.bru\\_prediction\(\)](#), [gg.matrix\(\)](#)

## Examples

```

if (bru_safe_inla() &&
    require(sn, quietly = TRUE) &&
    require(ggplot2, quietly = TRUE)) {
  # Generate some data

  input.df <- data.frame(x = cos(1:10))
  input.df <- within(input.df, y <- 5 + 2 * cos(1:10) + rnorm(10, mean = 0, sd = 0.1))

  # Fit a model with fixed effect 'x' and intercept 'Intercept'

  fit <- bru(y ~ x, family = "gaussian", data = input.df)

  # Predict posterior statistics of 'x'

  xpost <- predict(fit, NULL, formula = ~x_latent)

  # The statistics include mean, standard deviation, the 2.5% quantile, the median,
  # the 97.5% quantile, minimum and maximum sample drawn from the posterior as well as
  # the coefficient of variation and the variance.

  xpost

  # For a single variable like 'x' the default plotting method invoked by gg() will
  # show these statistics in a fashion similar to a box plot:
  ggplot() +
    gg(xpost)

  # The predict function can also be used to simultaneously estimate posteriors
  # of multiple variables:

  xipost <- predict(fit,
    newdata = NULL,
    formula = ~ c(
      Intercept = Intercept_latent,
      x = x_latent
    )
  )
  xipost

  # If we still want a plot in the previous style we have to set the bar parameter to TRUE

  p1 <- ggplot() +
    gg(xipost, bar = TRUE)
  p1

  # Note that gg also understands the posterior estimates generated while running INLA

  p2 <- ggplot() +
    gg(fit$summary.fixed, bar = TRUE)
  multiplot(p1, p2)

```

```

# By default, if the prediction has more than one row, gg will plot the column 'mean' against
# the row index. This is for instance useful for predicting and plotting function
# but not very meaningful given the above example:

ggplot() +
  gg(xipost)

# For ease of use we can also type

plot(xipost)

# This type of plot will show a ribbon around the mean, which visualizes the upper and lower
# quantiles mentioned above (2.5 and 97.5%). Plotting the ribbon can be turned off using the
# \code{ribbon} parameter

ggplot() +
  gg(xipost, ribbon = FALSE)

# Much like the other geoms produced by gg we can adjust the plot using ggplot2 style
# commands, for instance

ggplot() +
  gg(xipost) +
  gg(xipost, mapping = aes(y = median), ribbon = FALSE, color = "red")
}

```

**gg.fm\_mesh\_1d***Geom for fm\_mesh\_1d objects***Description**

This function generates a geom\_point object showing the knots (vertices) of a 1D mesh. Requires the ggplot2 package.

**Usage**

```

## S3 method for class 'fm_mesh_1d'
gg(
  data,
  mapping = ggplot2::aes(.data[["x"]], .data[["y"]]),
  y = 0,
  shape = 4,
  ...
)

```

### Arguments

<code>data</code>	An <code>fmesher::fm_mesh_1d</code> object.
<code>mapping</code>	aesthetic mappings created by <code>aes</code> . These are passed on to <code>geom_point</code> .
<code>y</code>	Single or vector numeric defining the y-coordinates of the mesh knots to plot.
<code>shape</code>	Shape of the knot markers.
<code>...</code>	parameters passed on to <code>geom_point</code> .

### Value

An object generated by `geom_point`.

### See Also

Other geomes for meshes: `gg()`, `gg.fm_mesh_2d()`

### Examples

```
if (require("fmesher", quietly = TRUE) &&
    require("ggplot2", quietly = TRUE)) {
  # Create a 1D mesh

  mesh <- fm_mesh_1d(seq(0, 10, by = 0.5))

  # Plot it

  ggplot() +
    gg(mesh)

  # Plot it using a different shape and size for the mesh nodes

  ggplot() +
    gg(mesh, shape = "|", size = 5)
}
```

`gg.fm_mesh_2d`      *Geom for fm\_mesh\_2d objects*

### Description

This function extracts the graph of an `fmesher::fm_mesh_2d` object and uses `geom_line` to visualize the graph's edges. Alternatively, if the `color` argument is provided, interpolates the colors across for a set of `SpatialPixels` covering the mesh area and calls `gg.SpatialPixelsDataFrame()` to plot the interpolation. Requires the `ggplot2` package.

Also see the `fmesher::geom_fm()` method.

**Usage**

```
## S3 method for class 'fm_mesh_2d'
gg(
  data,
  color = NULL,
  alpha = NULL,
  edge.color = "grey",
  edge.linewidth = 0.25,
  interior = TRUE,
  int.color = "blue",
  int.linewidth = 0.5,
  exterior = TRUE,
  ext.color = "black",
  ext.linewidth = 1,
  crs = NULL,
  mask = NULL,
  nx = 500,
  ny = 500,
  ...
)
```

**Arguments**

<code>data</code>	An <code>fm_mesh_2d</code> object.
<code>color</code>	A vector of scalar values to fill the mesh with colors. The length of the vector must correspond to the number of mesh vertices. The alternative name <code>colour</code> is also recognised.
<code>alpha</code>	A vector of scalar values setting the alpha value of the colors provided.
<code>edge.color</code>	Color of the regular mesh edges.
<code>edge.linewidth</code>	Line width for the regular mesh edges. Default 0.25
<code>interior</code>	If <code>TRUE</code> , plot the interior boundaries of the mesh.
<code>int.color</code>	Color used to plot the interior constraint edges.
<code>int.linewidth</code>	Line width for the interior constraint edges. Default 0.5
<code>exterior</code>	If <code>TRUE</code> , plot the exterior boundaries of the mesh.
<code>ext.color</code>	Color used to plot the exterior boundary edges.
<code>ext.linewidth</code>	Line width for the exterior boundary edges. Default 1
<code>crs</code>	A CRS object supported by <code>fmesher::fm_transform()</code> defining the coordinate system to project the mesh to before plotting.
<code>mask</code>	A <code>SpatialPolygon</code> or <code>sf</code> polygon defining the region that is plotted.
<code>nx</code>	Number of pixels in x direction (when plotting using the <code>color</code> parameter).
<code>ny</code>	Number of pixels in y direction (when plotting using the <code>color</code> parameter).
<code>...</code>	ignored arguments (S3 generic compatibility).

**Value**

`geom_line` return values or, if the color argument is used, the values of [gg.SpatialPixelsDataFrame\(\)](#).

**See Also**

Other geomes for meshes: [gg\(\)](#), [gg.fm\\_mesh\\_1d\(\)](#)

**Examples**

```
if (require(fmesher, quietly = TRUE) &&
    require(ggplot2, quietly = TRUE)) {

  # Load Gorilla data
  gorillas <- inlabru::gorillas_sf

  # Plot mesh using default edge colors

  ggplot() +
    gg(gorillas$mesh)

  # Don't show interior and exterior boundaries

  ggplot() +
    gg(gorillas$mesh, interior = FALSE, exterior = FALSE)

  # Change the edge colors

  ggplot() +
    gg(gorillas$mesh,
      edge.color = "green",
      int.color = "black",
      ext.color = "blue"
    )

  # Use the x-coordinate of the vertices to colorize the triangles and
  # mask the plotted area by the survey boundary, i.e. only plot the inside

  xcoord <- gorillas$mesh$loc[, 1]
  ggplot() +
    gg(gorillas$mesh, color = (xcoord - 580), mask = gorillas$boundary) +
    gg(gorillas$boundary, alpha = 0)
}
```

**Description**

Creates a tile geom for plotting a matrix

**Usage**

```
## S3 method for class 'matrix'  
gg(data, mapping = NULL, ...)
```

**Arguments**

data            A matrix object.  
mapping        a set of aesthetic mappings created by aes. These are passed on to geom\_tile.  
...              Arguments passed on to geom\_tile.

**Details**

Requires the ggplot2 package.

**Value**

A geom\_tile with reversed y scale.

**See Also**

Other geomes for inla and inlabru predictions: [gg\(\)](#), [gg.bru\\_prediction\(\)](#), [gg.data.frame\(\)](#)

**Examples**

```
if (require("ggplot2", quietly = TRUE)) {  
  A <- matrix(runif(100), nrow = 10)  
  ggplot() +  
    gg(A)  
}
```

---

gg.RasterLayer            *Geom for RasterLayer objects*

---

**Description**

This function takes a RasterLayer object, converts it into a SpatialPixelsDataFrame and uses geom\_tile to plot the data.

**Usage**

```
## S3 method for class 'RasterLayer'  
gg(  
  data,  
  mapping = ggplot2::aes(x = .data[["x"]], y = .data[["y"]], fill = .data[["layer"]]),  
  ...  
)
```

## Arguments

- `data` A RasterLayer object.
- `mapping` aesthetic mappings created by `aes`. These are passed on to `geom_tile`.
- `...` Arguments passed on to `geom_tile`.

## Details

This function requires the `raster` and `ggplot2` packages.

## Value

An object returned by `geom_tile`

## See Also

Other geomes for Raster data: [gg\(\)](#)

## Examples

```
## Not run:
# Some features require the raster and spatstat.data packages.
if (require("spatstat.data", quietly = TRUE) &&
    require("raster", quietly = TRUE) &&
    require("ggplot2", quietly = TRUE)) {
  # Load Gorilla data
  data("gorillas", package = "spatstat.data", envir = environment())

  # Convert elevation covariate to RasterLayer

  elev <- as(gorillas.extra$elevation, "RasterLayer")

  # Plot the elevation

  ggplot() +
    gg(elev)
}

## End(Not run)
```

## Description

This function uses `geom_sf()`, unless overridden by the `geom` argument. Requires the `ggplot2` package.

## Usage

```
## S3 method for class 'sf'  
gg(data, mapping = NULL, ..., geom = "sf")
```

## Arguments

data	An sf object.
mapping	Default mapping is ggplot2::aes(geometry = ...), where the geometry name is obtained from attr(data, "sf_column"). This is merged with the user supplied mapping.
...	Arguments passed on to geom_sf or geom_tile.
geom	Either "sf" (default) or "tile". For "tile", uses geom_tile(..., stat = "sf_coordinates"), intended for converting point data to grid tiles with the fill aesthetic, which is by default set to the first data column.

## Value

A ggplot return value

## See Also

Other geomes for spatial data: [gg\(\)](#), [gg.SpatRaster\(\)](#), [gg.SpatialGridDataFrame\(\)](#), [gg.SpatialLines\(\)](#), [gg.SpatialPixels\(\)](#), [gg.SpatialPixelsDataFrame\(\)](#), [gg.SpatialPoints\(\)](#), [gg.SpatialPolygons\(\)](#)

## Examples

```
if (require("ggplot2", quietly = TRUE) &&  
    requireNamespace("terra", quietly = TRUE) &&  
    require("tidyterra", quietly = TRUE)) {  
  # Load Gorilla data  
  
  gorillas <- inlabru::gorillas_sf  
  gorillas$gcov <- gorillas_sf_gcov()  
  
  # Plot Gorilla elevation covariate provided as terra::rast.  
  
  ggplot() +  
    gg(gorillas$gcov$elevation)  
  
  # Add Gorilla survey boundary and nest sightings  
  
  ggplot() +  
    gg(gorillas$gcov$elevation) +  
    gg(gorillas$boundary, alpha = 0) +  
    gg(gorillas$nests)  
  
  # Load pantropical dolphin data  
  
  mexdolphin <- inlabru::mexdolphin_sf
```

```

# Plot the pantropical survey boundary, ship transects and dolphin sightings

ggplot() +
  gg(mexdolphin$ppoly, alpha = 0.5) + # survey boundary
  gg(mexdolphin$samplers) + # ship transects
  gg(mexdolphin$points) # dolphin sightings

# Change color

ggplot() +
  gg(mexdolphin$ppoly, color = "green", alpha = 0.5) + # survey boundary
  gg(mexdolphin$samplers, color = "red") + # ship transects
  gg(mexdolphin$points, color = "blue") # dolphin sightings

# Visualize data annotations: line width by segment number

names(mexdolphin$samplers) # 'seg' holds the segment number
ggplot() +
  gg(mexdolphin$samplers, aes(color = seg))

# Visualize data annotations: point size by dolphin group size

names(mexdolphin$points) # 'size' holds the group size
ggplot() +
  gg(mexdolphin$points, aes(size = size))
}

```

**gg.SpatialGridDataFrame***Geom for SpatialGridDataFrame objects***Description**

Coerces input `SpatialGridDataFrame` to `SpatialPixelsDataFrame` and calls [gg.SpatialPixelsDataFrame\(\)](#) to plot it. Requires the `ggplot2` package.

**Usage**

```
## S3 method for class 'SpatialGridDataFrame'
gg(data, ...)
```

**Arguments**

- |                   |  |
|-------------------|--|
| <code>data</code> | A <code>SpatialGridDataFrame</code> object.                          |
| <code>...</code>  | Arguments passed on to <a href="#">gg.SpatialPixelsDataFrame()</a> . |

**Value**

A geom\_tile value.

**See Also**

Other geomes for spatial data: [gg\(\)](#), [gg.SpatRaster\(\)](#), [gg.SpatialLines\(\)](#), [gg.SpatialPixels\(\)](#), [gg.SpatialPixelsDataFrame\(\)](#), [gg.SpatialPoints\(\)](#), [gg.SpatialPolygons\(\)](#), [gg.sf\(\)](#)

**Examples**

```
if (require("ggplot2", quietly = TRUE) &&
    requireNamespace("terra", quietly = TRUE) &&
    bru_safe_sp() &&
    require("sp")) {
  # Load Gorilla data

  gorillas <- inlabru::gorillas_sf

  gcov <- gorillas_sf_gcov()
  elev <- terra::as.data.frame(gcov$elevation, xy = TRUE)
  elev <- sf::as_Spatial(sf::st_as_sf(elev, coords = c("x", "y")))

  # Turn elevation covariate into SpatialGridDataFrame
  elev <- sp::SpatialPixelsDataFrame(elev, data = as.data.frame(elev))

  # Plot Gorilla elevation covariate provided as SpatialPixelsDataFrame.
  # The same syntax applies to SpatialGridDataFrame objects.

  ggplot() +
    gg(elev)

  # Add Gorilla survey boundary and nest sightings

  ggplot() +
    gg(elev) +
    gg(gorillas$boundary, alpha = 0.0, col = "red") +
    gg(gorillas$nests)

  # Load pantropical dolphin data

  mexdolphin <- inlabru::mexdolphin_sp()

  # Plot the pantropical survey boundary, ship transects and dolphin sightings

  ggplot() +
    gg(mxdolphin$ppoly) + # survey boundary as SpatialPolygon
    gg(mxdolphin$samplers) + # ship transects as SpatialLines
    gg(mxdolphin$points) # dolphin sightings as SpatialPoints

  # Change color

  ggplot() +
```

```

gg(mexdolphin$ppoly, color = "green") + # survey boundary as SpatialPolygon
gg(mexdolphin$samplers, color = "red") + # ship transects as SpatialLines
gg(mexdolphin$points, color = "blue") # dolphin sightings as SpatialPoints

# Visualize data annotations: line width by segment number

names(mexdolphin$samplers) # 'seg' holds the segment number
ggplot() +
  gg(mexdolphin$samplers, aes(color = seg))

# Visualize data annotations: point size by dolphin group size

names(mexdolphin$points) # 'size' holds the group size
ggplot() +
  gg(mexdolphin$points, aes(size = size))
}

```

**gg.SpatialLines** *Geom for SpatialLines objects*

## Description

Extracts start and end points of the lines and calls `geom_segment` to plot lines between them. Requires the `ggplot2` package.

## Usage

```
## S3 method for class 'SpatialLines'
gg(data, mapping = NULL, crs = NULL, ...)
```

## Arguments

- |                      |  |
|----------------------|--|
| <code>data</code>    | A <code>SpatialLines</code> or <code>SpatialLinesDataFrame</code> object.  |
| <code>mapping</code> | Aesthetic mappings created by <code>ggplot2::aes</code> or <code>ggplot2::aes_</code> used to update the default mapping. The default mapping is<br><br><code>ggplot2::aes(</code><br><code>  x = .data[[sp::coordnames(data)[1]]],</code><br><code>  y = .data[[sp::coordnames(data)[2]]],</code><br><code>  xend = .data[[paste0("end.", sp::coordnames(data)[1])]],</code><br><code>  yend = .data[[paste0("end.", sp::coordnames(data)[2])]])</code> |
| <code>crs</code>     | A CRS object defining the coordinate system to project the data to before plotting.  |
| ...                  | Arguments passed on to <code>ggplot2::geom_segment</code> .  |

## Value

A ‘geom\_segment’ return value.

## See Also

Other geomes for spatial data: [gg\(\)](#), [gg.SpatRaster\(\)](#), [gg.SpatialGridDataFrame\(\)](#), [gg.SpatialPixels\(\)](#), [gg.SpatialPixelsDataFrame\(\)](#), [gg.SpatialPoints\(\)](#), [gg.SpatialPolygons\(\)](#), [gg.sf\(\)](#)

## Examples

```
if (require("ggplot2", quietly = TRUE) &&
    requireNamespace("terra", quietly = TRUE) &&
    bru_safe_sp() &&
    require("sp")) {
  # Load Gorilla data

  gorillas <- inlabru::gorillas_sf

  gcov <- gorillas_sf_gcov()
  elev <- terra::as.data.frame(gcov$elevation, xy = TRUE)
  elev <- sf::as_Spatial(sf::st_as_sf(elev, coords = c("x", "y")))

  # Turn elevation covariate into SpatialGridDataFrame
  elev <- sp::SpatialPixelsDataFrame(elev, data = as.data.frame(elev))

  # Plot Gorilla elevation covariate provided as SpatialPixelsDataFrame.
  # The same syntax applies to SpatialGridDataFrame objects.

  ggplot() +
    gg(elev)

  # Add Gorilla survey boundary and nest sightings

  ggplot() +
    gg(elev) +
    gg(gorillas$boundary, alpha = 0.0, col = "red") +
    gg(gorillas$nests)

  # Load pantropical dolphin data

  mexdolphin <- inlabru::mexdolphin_sp()

  # Plot the pantropical survey boundary, ship transects and dolphin sightings

  ggplot() +
    gg(mexdolphin$poly) + # survey boundary as SpatialPolygon
    gg(mexdolphin$samplers) + # ship transects as SpatialLines
    gg(mexdolphin$points) # dolphin sightings as SpatialPoints

  # Change color

  ggplot() +
    gg(mexdolphin$poly, color = "green") + # survey boundary as SpatialPolygon
    gg(mexdolphin$samplers, color = "red") + # ship transects as SpatialLines
    gg(mexdolphin$points, color = "blue") # dolphin sightings as SpatialPoints
```

```

# Visualize data annotations: line width by segment number

names(mexdolphin$samplers) # 'seg' holds the segment number
ggplot() +
  gg(mexdolphin$samplers, aes(color = seg))

# Visualize data annotations: point size by dolphin group size

names(mexdolphin$points) # 'size' holds the group size
ggplot() +
  gg(mexdolphin$points, aes(size = size))
}

```

**gg.SpatialPixels**      *Geom for SpatialPixels objects*

## Description

Uses `geom_point` to plot the pixel centers. Requires the `ggplot2` package.

## Usage

```
## S3 method for class 'SpatialPixels'
gg(data, ...)
```

## Arguments

data	A <code>sp::SpatialPixels</code> object.
...	Arguments passed on to <code>geom_tile</code> .

## Value

A `geom_tile` return value.

## See Also

Other geomes for spatial data: `gg()`, `gg.SpatRaster()`, `gg.SpatialGridDataFrame()`, `gg.SpatialLines()`, `gg.SpatialPixelsDataFrame()`, `gg.SpatialPoints()`, `gg.SpatialPolygons()`, `gg.sf()`

## Examples

```

if (require("ggplot2", quietly = TRUE) &&
  requireNamespace("terra", quietly = TRUE) &&
  bru_safe_sp()) {
  # Load Gorilla data

  gcov <- gorillas_sf_gcov()
```

```

elev <- terra::as.data.frame(gcov$elevation, xy = TRUE)
pxl <- sf:::as_Spatial(sf:::st_as_sf(elev, coords = c("x", "y")))

# Turn elevation covariate into SpatialPixels
pxl <- sp:::SpatialPixels(pxl)

# Plot the pixel centers
ggplot() +
  gg(px1, size = 0.1)
}

```

**gg.SpatialPixelsDataFrame***Geom for SpatialPixelsDataFrame objects***Description**

Coerces input SpatialPixelsDataFrame to data.frame and uses geom\_tile to plot it. Requires the ggplot2 package.

**Usage**

```
## S3 method for class 'SpatialPixelsDataFrame'
gg(data, mapping = NULL, crs = NULL, mask = NULL, ...)
```

**Arguments**

- |   |  |
|---|--|
| data  | A SpatialPixelsDataFrame object.   |
| mapping   | Aesthetic mappings created by aes used to update the default mapping. The default mapping is |
| <pre>ggplot2::aes(   x = .data[[sp::coordnames(data)[1]]],   y = .data[[sp::coordnames(data)[2]]],   fill = .data[[names(data)[1]]] )</pre> |  |
| crs   | A sp::CRS object defining the coordinate system to project the data to before plotting.      |
| mask  | A sp::SpatialPolygons object defining the region that is plotted.                            |
| ...   | Arguments passed on to geom_tile.  |

**Value**

A geom\_tile return value.

**See Also**

Other geomes for spatial data: [gg\(\)](#), [gg.SpatRaster\(\)](#), [gg.SpatialGridDataFrame\(\)](#), [gg.SpatialLines\(\)](#), [gg.SpatialPixels\(\)](#), [gg.SpatialPoints\(\)](#), [gg.SpatialPolygons\(\)](#), [gg.sf\(\)](#)

**Examples**

```
if (require("ggplot2", quietly = TRUE) &&
    requireNamespace("terra", quietly = TRUE) &&
    bru_safe_sp() &&
    require("sp")) {
  # Load Gorilla data

  gorillas <- inlabru::gorillas_sf

  gcov <- gorillas_sf_gcov()
  elev <- terra::as.data.frame(gcov$elevation, xy = TRUE)
  elev <- sf:::as_Spatial(sf:::st_as_sf(elev, coords = c("x", "y")))

  # Turn elevation covariate into SpatialGridDataFrame
  elev <- sp:::SpatialPixelsDataFrame(elev, data = as.data.frame(elev))

  # Plot Gorilla elevation covariate provided as SpatialPixelsDataFrame.
  # The same syntax applies to SpatialGridDataFrame objects.

  ggplot() +
    gg(elev)

  # Add Gorilla survey boundary and nest sightings

  ggplot() +
    gg(elev) +
    gg(gorillas$boundary, alpha = 0.0, col = "red") +
    gg(gorillas$nests)

  # Load pantropical dolphin data

  mexdolphin <- inlabru::mexdolphin_sp()

  # Plot the pantropical survey boundary, ship transects and dolphin sightings

  ggplot() +
    gg(mxdolphin$ppoly) + # survey boundary as SpatialPolygon
    gg(mxdolphin$samplers) + # ship transects as SpatialLines
    gg(mxdolphin$points) # dolphin sightings as SpatialPoints

  # Change color

  ggplot() +
    gg(mxdolphin$ppoly, color = "green") + # survey boundary as SpatialPolygon
    gg(mxdolphin$samplers, color = "red") + # ship transects as SpatialLines
    gg(mxdolphin$points, color = "blue") # dolphin sightings as SpatialPoints
```

```
# Visualize data annotations: line width by segment number  
  
names(mexdolphin$samplers) # 'seg' holds the segment number  
ggplot() +  
  gg(mexdolphin$samplers, aes(color = seg))  
  
# Visualize data annotations: point size by dolphin group size  
  
names(mexdolphin$points) # 'size' holds the group size  
ggplot() +  
  gg(mexdolphin$points, aes(size = size))  
}
```

---

gg.SpatialPoints      *Geom for SpatialPoints objects*

---

## Description

This function coerces the SpatialPoints into a data.frame and uses geom\_point to plot the points. Requires the ggplot2 package.

## Usage

```
## S3 method for class 'SpatialPoints'  
gg(data, mapping = NULL, crs = NULL, ...)
```

## Arguments

- |         |   |
|---------|---|
| data    | A SpatialPoints object.   |
| mapping | Aesthetic mappings created by aes used to update the default mapping. The default mapping is<br><br>ggplot2::aes(<br>x = .data[[sp::coordnames(data)[1]]],<br>y = .data[[sp::coordnames(data)[2]]]<br>) |
| crs     | A sp::CRS object defining the coordinate system to project the data to before plotting.   |
| ...     | Arguments passed on to geom_point.  |

## Value

A geom\_point return value

**See Also**

Other geomes for spatial data: [gg\(\)](#), [gg.SpatRaster\(\)](#), [gg.SpatialGridDataFrame\(\)](#), [gg.SpatialLines\(\)](#), [gg.SpatialPixels\(\)](#), [gg.SpatialPixelsDataFrame\(\)](#), [gg.SpatialPolygons\(\)](#), [gg.sf\(\)](#)

**Examples**

```
if (require("ggplot2", quietly = TRUE) &&
    requireNamespace("terra", quietly = TRUE) &&
    bru_safe_sp() &&
    require("sp")) {
  # Load Gorilla data

  gorillas <- inlabru::gorillas_sf

  gcov <- gorillas_sf_gcov()
  elev <- terra::as.data.frame(gcov$elevation, xy = TRUE)
  elev <- sf:::as_Spatial(sf:::st_as_sf(elev, coords = c("x", "y")))

  # Turn elevation covariate into SpatialGridDataFrame
  elev <- sp:::SpatialPixelsDataFrame(elev, data = as.data.frame(elev))

  # Plot Gorilla elevation covariate provided as SpatialPixelsDataFrame.
  # The same syntax applies to SpatialGridDataFrame objects.

  ggplot() +
    gg(elev)

  # Add Gorilla survey boundary and nest sightings

  ggplot() +
    gg(elev) +
    gg(gorillas$boundary, alpha = 0.0, col = "red") +
    gg(gorillas$nests)

  # Load pantropical dolphin data

  mexdolphin <- inlabru::mexdolphin_sp()

  # Plot the pantropical survey boundary, ship transects and dolphin sightings

  ggplot() +
    gg(mxdolphin$ppoly) + # survey boundary as SpatialPolygon
    gg(mxdolphin$samplers) + # ship transects as SpatialLines
    gg(mxdolphin$points) # dolphin sightings as SpatialPoints

  # Change color

  ggplot() +
    gg(mxdolphin$ppoly, color = "green") + # survey boundary as SpatialPolygon
    gg(mxdolphin$samplers, color = "red") + # ship transects as SpatialLines
    gg(mxdolphin$points, color = "blue") # dolphin sightings as SpatialPoints
```

```

# Visualize data annotations: line width by segment number

names(mexdolphin$samplers) # 'seg' holds the segment number
ggplot() +
  gg(mexdolphin$samplers, aes(color = seg))

# Visualize data annotations: point size by dolphin group size

names(mexdolphin$points) # 'size' holds the group size
ggplot() +
  gg(mexdolphin$points, aes(size = size))
}

```

`gg.SpatialPolygons`      *Geom for SpatialPolygons objects*

## Description

Uses the `ggplot2::fortify()` function to turn the `SpatialPolygons` objects into a `data.frame`. Then calls `geom_polygon` to plot the polygons. Requires the `ggplot2` package.

## Usage

```
## S3 method for class 'SpatialPolygons'
gg(data, mapping = NULL, crs = NULL, ...)
```

## Arguments

<code>data</code>	A <code>SpatialPolygons</code> or <code>SpatialPolygonsDataFrame</code> object.
<code>mapping</code>	Aesthetic mappings created by <code>aes</code> used to update the default mapping.
<code>crs</code>	A CRS object defining the coordinate system to project the data to before plotting.
<code>...</code>	Arguments passed on to <code>geom_sf</code> . Unless specified by the user, the argument <code>alpha = 0.2</code> (alpha level for polygon filling) is added.

## Details

Up to version 2.10.0, the `ggeopath` package was used to ensure proper plotting, since the `ggplot2::geom_polygon` function doesn't always handle geometries with holes properly. After 2.10.0, the object is converted to `sf` format and passed on to `gg.sf()` instead, as `ggplot2` version 3.4.4 deprecated the internally used `ggplot2::fortify()` method for `SpatialPolygons/DataFrame` objects.

## Value

A `geom_sf` object.

**See Also**

Other geomes for spatial data: [gg\(\)](#), [gg.SpatRaster\(\)](#), [gg.SpatialGridDataFrame\(\)](#), [gg.SpatialLines\(\)](#), [gg.SpatialPixels\(\)](#), [gg.SpatialPixelsDataFrame\(\)](#), [gg.SpatialPoints\(\)](#), [gg.sf\(\)](#)

**Examples**

```
if (require("ggplot2", quietly = TRUE) &&
    requireNamespace("terra", quietly = TRUE) &&
    bru_safe_sp() &&
    require("sp")) {
  # Load Gorilla data

  gorillas <- inlabru::gorillas_sf

  gcov <- gorillas_sf_gcov()
  elev <- terra::as.data.frame(gcov$elevation, xy = TRUE)
  elev <- sf:::as_Spatial(sf:::st_as_sf(elev, coords = c("x", "y")))

  # Turn elevation covariate into SpatialGridDataFrame
  elev <- sp:::SpatialPixelsDataFrame(elev, data = as.data.frame(elev))

  # Plot Gorilla elevation covariate provided as SpatialPixelsDataFrame.
  # The same syntax applies to SpatialGridDataFrame objects.

  ggplot() +
    gg(elev)

  # Add Gorilla survey boundary and nest sightings

  ggplot() +
    gg(elev) +
    gg(gorillas$boundary, alpha = 0.0, col = "red") +
    gg(gorillas$nests)

  # Load pantropical dolphin data

  mexdolphin <- inlabru::mexdolphin_sp()

  # Plot the pantropical survey boundary, ship transects and dolphin sightings

  ggplot() +
    gg(mexdolphin$poly) + # survey boundary as SpatialPolygon
    gg(mexdolphin$samplers) + # ship transects as SpatialLines
    gg(mexdolphin$points) # dolphin sightings as SpatialPoints

  # Change color

  ggplot() +
    gg(mexdolphin$poly, color = "green") + # survey boundary as SpatialPolygon
    gg(mexdolphin$samplers, color = "red") + # ship transects as SpatialLines
    gg(mexdolphin$points, color = "blue") # dolphin sightings as SpatialPoints
```

```
# Visualize data annotations: line width by segment number  
  
names(mexdolphin$samplers) # 'seg' holds the segment number  
ggplot() +  
  gg(mexdolphin$samplers, aes(color = seg))  
  
# Visualize data annotations: point size by dolphin group size  
  
names(mexdolphin$points) # 'size' holds the group size  
ggplot() +  
  gg(mexdolphin$points, aes(size = size))  
}
```

---

**gg.SpatRaster**

*Geom wrapper for SpatRaster objects*

---

**Description**

Convenience wrapper function for `tidyterra::geom_spatraster()`. Requires the `ggplot2` and `tidyterra` packages.

**Usage**

```
## S3 method for class 'SpatRaster'  
gg(data, ...)
```

**Arguments**

`data` A SpatRaster object.  
`...` Arguments passed on to `geom_spatraster`.

**Value**

The output from ‘`geom_spatraster`’.

**See Also**

Other geomes for spatial data: `gg()`, `gg.SpatialGridDataFrame()`, `gg.SpatialLines()`, `gg.SpatialPixels()`, `gg.SpatialPixelsDataFrame()`, `gg.SpatialPoints()`, `gg.SpatialPolygons()`, `gg.sf()`

**Examples**

```
if (require("ggplot2", quietly = TRUE) &&  
  requireNamespace("terra", quietly = TRUE) &&  
  require("tidyterra", quietly = TRUE)) {  
  # Load Gorilla covariates
```

```
gcov <- gorillas_sf_gcov()

# Plot the pixel centers
ggplot() +
  gg(gcov$elevation)
}
```

**globe** *Visualize a globe using RGL*

## Description

Creates a textured sphere and lon/lat coordinate annotations. This function requires the `rgl` and `sphereplot` packages.

## Usage

```
globe(
  R = 1,
  R.grid = 1.05,
  specular = "black",
  axes = FALSE,
  box = FALSE,
  xlab = "",
  ylab = "",
  zlab = "")
```

## Arguments

R	Radius of the globe
R.grid	Radius of the annotation sphere.
specular	Light color of specular effect.
axes	If TRUE, plot x, y and z axes.
box	If TRUE, plot a box around the globe.
xlab, ylab, zlab	Axes labels

## Value

No value, used for plotting side effect.

## See Also

Other inlabru RGL tools: [glplot\(\)](#)

## Examples

```

if (interactive() &&
    require("rgl", quietly = TRUE) &&
    require("sphereplot", quietly = TRUE) &&
    bru_safe_sp() &&
    require("sp")) {
  # Show the globe
  globe()

  # Load pantropical dolphin data
  mexdolphin <- inlabru::mexdolphin_sp()

  # Add mesh, ship transects and dolphin sightings stored
  # as fm_mesh_2d, SpatialLines and SpatialPoints objects, respectively

  glplot(mxdolphin$mesh, alpha = 0.2)
  glplot(mxdolphin$samplers, lwd = 5)
  glplot(mxdolphin$points, size = 10)
}

```

## glplot

*Render objects using RGL*

## Description

glplot() is a generic function for renders various kinds of spatial objects, i.e. Spatial\* data and fm\_mesh\_2d objects. The function invokes particular methods which depend on the class of the first argument.

## Usage

```

glplot(object, ...)

## S3 method for class 'SpatialPoints'
glplot(object, add = TRUE, color = "red", ...)

## S3 method for class 'SpatialLines'
glplot(object, add = TRUE, ...)

## S3 method for class 'fm_mesh_2d'
glplot(object, add = TRUE, col = NULL, ...)

```

## Arguments

object	an object used to select a method.
...	Parameters passed on to plot_rgl.fm_mesh_2d()
add	If TRUE, add the points to an existing plot. If FALSE, create new plot.

<code>color</code>	vector of R color characters. See <code>material3d()</code> for details.
<code>col</code>	Color specification. A single named color, a vector of scalar values, or a matrix of RGB values.

### Methods (by class)

- `glplot(SpatialPoints)`: This function will calculate the cartesian coordinates of the points provided and use `points3d()` in order to render them.
- `glplot(SpatialLines)`: This function will calculate a cartesian representation of the lines provided and use `lines3d()` in order to render them.
- `glplot(fm_mesh_2d)`: This function transforms the mesh to 3D cartesian coordinates and uses `fmesher::plot_rgl()` to plot the result.

### See Also

Other inlabru RGL tools: [globe\(\)](#)

### Examples

```
if (interactive() &&
    require("rgl", quietly = TRUE) &&
    require("sphereplot", quietly = TRUE) &&
    bru_safe_sp() &&
    require("sp")) {
  # Show the globe
  globe()

  # Load pantropical dolphin data
  mexdolphin <- inlabru::mexdolphin_sp()

  # Add mesh, ship transects and dolphin sightings stored
  # as fm_mesh_2d, SpatialLines and SpatialPoints objects, respectively

  glplot(mexdolphin$mesh, alpha = 0.2)
  glplot(mexdolphin$samplers, lwd = 5)
  glplot(mexdolphin$points, size = 10)
}
```

---

<code>gorillas_sf</code>	<i>Gorilla nesting sites in sf format</i>
--------------------------	---

---

## Description

This is the gorillas dataset from the package `spatstat.data`, reformatted as point process data for use with `inlabru`.

## Usage

```
gorillas_sf
data(gorillas_sf, package = "inlabru")

gorillas_sf_gcov()

gorillas_sp()
```

## Format

The data are a list that contains these elements:

**nests:** An `sf` object containing the locations of the gorilla nests.

**boundary:** An `sf` object defining the boundary of the region that was searched for the nests.

**mesh:** An `fm_mesh_2d` object containing a mesh that can be used with function `lgcp` to fit a LGCP to the nest data.

**gcov\_file:** The in-package filename of a `terra::SpatRaster` object, with one layer for each of these spatial covariates:

- aspect** Compass direction of the terrain slope. Categorical, with levels N, NE, E, SE, S, SW, W and NW, which are coded as integers 1 to 8.
- elevation** Digital elevation of terrain, in metres.
- heat** Heat Load Index at each point on the surface (Beer's aspect), discretised. Categorical with values Warmest (Beer's aspect between 0 and 0.999), Moderate (Beer's aspect between 1 and 1.999), Coolest (Beer's aspect equals 2). These are coded as integers 1, 2 and 3, in that order.
- slopangle** Terrain slope, in degrees.
- slopetype** Type of slope. Categorical, with values Valley, Toe (toe slope), Flat, Midslope, Upper and Ridge. These are coded as integers 1 to 6.
- vegetation** Vegetation type: a categorical variable with 6 levels coded as integers 1 to 6 (in order of increasing expected habitat suitability)
- waterdist** Euclidean distance from nearest water body, in metres.

Loading of the covariates can be done with `gorillas_sf_gcov()` or

```
gorillas_sf$gcov <- terra::rast(
  system.file(gorillas_sf$gcov_file, package = "inlabru")
)
```

```

plotsample Plot sample of gorilla nests, sampling 9x9 over the region, with 60\
counts An sf object with elements count, exposure, and geometry, holding the point ge-\
ometry for the centre of each plot, the count in each plot and the area of each plot.
plots An sf object with MULTIPOLYGON objects defining the individual plot boundaries and
an all-ones weight column.
nests An sf giving the locations of each detected nests, group ("minor" or "major"), season
("dry" or "rainy"), and date (in Date format).

```

## Functions

- `gorillas_sf_gcov()`: Access the `gorillas_sf` covariates data as a `terra::rast()` object.
- `gorillas_sp()`: Access the `gorillas_sf` data in `sp` format. The covariate data is added as `gcov`, a list of `sp::SpatialPixelsDataFrame` objects. Requires the `sp`, `sf`, and `terra` packages to be installed.

## Source

Library `spatstat.data`.

## References

- Funwi-Gabga, N. (2008) A pastoralist survey and fire impact assessment in the Kagwene Gorilla Sanctuary, Cameroon. M.Sc. thesis, Geology and Environmental Science, University of Buea, Cameroon.
- Funwi-Gabga, N. and Mateu, J. (2012) Understanding the nesting spatial behaviour of gorillas in the Kagwene Sanctuary, Cameroon. Stochastic Environmental Research and Risk Assessment 26 (6), 793-811.

## Examples

```

if (interactive() &&
require(ggplot2, quietly = TRUE) &&
requireNamespace("terra", quietly = TRUE) &&
requireNamespace("tidyterra", quietly = TRUE)) {
  # plot all the nests, mesh and boundary
  ggplot() +
    gg(gorillas_sf$mesh) +
    geom_sf(
      data = gorillas_sf$boundary,
      alpha = 0.1, fill = "blue"
    ) +
    geom_sf(data = gorillas_sf$nests)

  # Plot the elevation covariate
  gorillas_sf$gcov <- gorillas_sf_gcov()
  ggplot() +
    tidyterra::geom_spatraster(data = gorillas_sf$gcov$elevation)

  # Plot the plot sample
  ggplot() +

```

```

    geom_sf(data = gorillas_sf$plotsample$plots) +
    geom_sf(data = gorillas_sf$plotsample$ests)
}
if (interactive() &&
  requireNamespace("terra", quietly = TRUE)) {
  gorillas_sf$gcov <- gorillas_sf_gcov()
}

```

lgcp

*Log Gaussian Cox process (LGCP) inference using INLA*

## Description

This function performs inference on a LGCP observed via points residing possibly multiple dimensions. These dimensions are defined via the left hand side of the formula provided via the model parameter. The left hand side determines the intensity function that is assumed to drive the LGCP. This may include effects that lead to a thinning (filtering) of the point process. By default, the log intensity is assumed to be a linear combination of the effects defined by the formula's RHS.

More sophisticated models, e.g. non-linear thinning, can be achieved by using the predictor argument. The latter requires multiple runs of INLA for improving the required approximation of the predictor. In many applications the LGCP is only observed through subsets of the dimensions the process is living in. For example, spatial point realizations may only be known in sub-areas of the modelled space. These observed subsets of the LGCP domain are called samplers and can be provided via the respective parameter. If samplers is NULL it is assumed that all of the LGCP's dimensions have been observed completely.

## Usage

```

lgcp(
  components,
  data,
  domain = NULL,
  samplers = NULL,
  ips = NULL,
  formula = . ~ .,
  ...,
  options = list(),
  .envir = parent.frame()
)

```

## Arguments

<code>components</code>	A formula-like specification of latent components. Also used to define a default linear additive predictor. See <a href="#">bru_comp()</a> for details.
<code>data</code>	Predictor expression-specific data, as a <code>data.frame</code> , <code>tibble</code> , or <code>sf</code> . Since 2.12.0.9023, deprecated support for <code>SpatialPoints[DataFrame]</code> objects.

domain, samplers, ips	Arguments used for family="cp" and aggregate=.
	domain Named list of domain definitions, see <a href="#">fmesher::fm_int()</a> .
	samplers Integration domain for family="cp" or subdomains for aggregate=, see <a href="#">fmesher::fm_int()</a> .
	ips Integration points. Defaults to fmesher::fm_int(domain, samplers). If explicitly given, overrides domain and samplers.
formula	a formula where the right hand side is a general R expression defines the predictor used in the model.
...	Further arguments passed on to <a href="#">bru_obs()</a> . In particular, optional E, a single numeric used rescale all integration weights by a fixed factor.
options	A <a href="#">bru_options</a> options object or a list of options passed on to <a href="#">bru_options()</a>
.envir	The evaluation environment to use for special arguments (E, Ntrials, weights, and scale) if not found in response_data or data. Defaults to the calling environment.

## Value

An [bru\(\)](#) object

## Examples

```
if (bru_safe_inla() &&
  require(ggplot2, quietly = TRUE) &&
  require(fmesher, quietly = TRUE) &&
  require(sn, quietly = TRUE)) {
  # Load the Gorilla data
  data <- gorillas_sf

  # Plot the Gorilla nests, the mesh and the survey boundary
  ggplot() +
    geom_fm(data = data$mesh) +
    gg(data$boundary, fill = "blue", alpha = 0.2) +
    gg(data$nests, col = "red", alpha = 0.2)

  # Define SPDE prior
  matern <- INLA::inla.spde2.pcmatern(
    data$mesh,
    prior.sigma = c(0.1, 0.01),
    prior.range = c(0.1, 0.01)
  )

  # Define domain of the LGCP as well as the model components (spatial SPDE
  # effect and Intercept)
  cmp <- geometry ~ field(geometry, model = matern) + Intercept(1)

  # Fit the model (with int.strategy="eb" to make the example take less time)
  fit <- lgcp(cmp, data$nests,
    samplers = data$boundary,
    domain = list(geometry = data$mesh),
```

```

    options = list(control.inla = list(int.strategy = "eb"))
  )

# Predict the spatial intensity surface
lambda <- predict(
  fit,
  fm_pixels(data$mesh, mask = data$boundary),
  ~ exp(field + Intercept)
)

# Plot the intensity
ggplot() +
  gg(lambda, geom = "tile") +
  geom_fm(data = data$mesh, alpha = 0, linewidth = 0.05) +
  gg(data$nests, col = "red", alpha = 0.2)
}

```

mexdolphin\_sf

*Pan-tropical spotted dolphins in the Gulf of Mexico*

## Description

This a version of the `mexdolphins` dataset from the package `dsm`, reformatted as point process data for use with `inlabru`, with the parts stored in `sf` format. The data are from a combination of several NOAA shipboard surveys conducted on pan-tropical spotted dolphins in the Gulf of Mexico. 47 observations of groups of dolphins were detected. The group size was recorded, as well as the Beaufort sea state at the time of the observation. Transect width is 16 km, i.e. maximal detection distance 8 km (transect half-width 8 km).

## Usage

```

mexdolphin_sf

mexdolphin_sp()

```

## Format

A list of objects:

- points:** An `sf` object containing the locations of detected dolphin groups, with their size as an attribute.
- samplers:** An `sf` object containing the transect lines that were surveyed.
- mesh:** An `fm_mesh_2d` object containing a Delaunay triangulation mesh (a type of discretization of continuous space) covering the survey region.
- ppoly:** An `sf` object defining the boundary of the survey region.
- simulated:** A `sf` object containing the locations of a *simulated* population of dolphin groups. The population was simulated from a `inlabru` model fitted to the actual survey data. Note that the simulated data do not have any associated size information.

## Functions

- `mxdolphin_sp()`: Convert `mxdolphin_sf` to `sp` format. Replaces the old `mxdolphin` dataset.

## Source

Library `dsm`.

## References

Halpin, P.N., A.J. Read, E. Fujioka, B.D. Best, B. Donnelly, L.J. Hazen, C. Kot, K. Urian, E. LaBrecque, A. Dimatteo, J. Cleary, C. Good, L.B. Crowder, and K.D. Hyrenbach. 2009. OBIS-SEAMAP: The world data center for marine mammal, sea bird, and sea turtle distributions. *Oceanography* 22(2):104-115

NOAA Southeast Fisheries Science Center. 1996. Report of a Cetacean Survey of Oceanic and Selected Continental Shelf Waters of the Northern Gulf of Mexico aboard NOAA Ship Oregon II (Cruise 220)

## Examples

```
if (require("ggplot2", quietly = TRUE)) {
  data(mxdolphin_sf, package = "inlabru", envir = environment())
  ggplot() +
    gg(mxdolphin_sf$mesh) +
    gg(mxdolphin_sf$ppoly, color = "blue", alpha = 0, linewidth = 1) +
    gg(mxdolphin_sf$samplers) +
    gg(mxdolphin_sf$points, aes(size = size), color = "red") +
    scale_size_area()

  ggplot() +
    gg(mxdolphin_sf$mesh,
       color = mxdolphin_sf$lambda,
       mask = mxdolphin_sf$ppoly
     )
}

if (require("ggplot2", quietly = TRUE) &&
  require("sp", quietly = TRUE)) {
  mxdolphin <- mxdolphin_sp()
  ggplot() +
    gg(mxdolphin$mesh) +
    gg(mxdolphin$ppoly, color = "blue") +
    gg(mxdolphin$samplers) +
    gg(mxdolphin$points, aes(size = size), color = "red") +
    scale_size_area() +
    coord_equal()

  ggplot() +
    gg(mxdolphin$mesh,
       col = mxdolphin$lambda,
```

```
    mask = mexdolphin$ppoly
) +
coord_equal()
}
```

## Description

Data imported from package MRSea, see <https://www.creem.st-andrews.ac.uk/software/>

## Usage

```
mrsea
```

## Format

A list of objects:

points A sf object containing the locations of XXXXX.  
samplers A sf object containing the transect lines that were surveyed.  
mesh An fm\_mesh\_2d object containing a Delaunay triangulation mesh (a type of discretization of continuous space) covering the survey region.  
boundary An sf object defining the boundary polygon of the survey region.  
covar An sf containing sea depth estimates.

## Source

Library MRSea.

## References

NONE YET

## Examples

```
if (require(ggplot2, quietly = TRUE)) {
  ggplot() +
    geom_fm(data = mrsea$mesh) +
    gg(mrsea$samplers) +
    gg(mrsea$points) +
    gg(mrsea$boundary)
}
```

<code>multiplot</code>	<i>Multiple ggplots on a page.</i>
------------------------	------------------------------------

## Description

[Deprecated] in favour of the `patchwork` package; see the example below.

Renders multiple ggplots on a single page.

## Usage

```
multiplot(..., plotlist = NULL, cols = 1, layout = NULL)
```

## Arguments

...	Comma-separated ggplot objects.
<code>plotlist</code>	A list of ggplot objects - an alternative to the comma-separated argument above.
<code>cols</code>	Number of columns of plots on the page.
<code>layout</code>	A matrix specifying the layout. If present, <code>cols</code> is ignored. If the layout is something like <code>matrix(c(1, 2, 3, 3), nrow=2, byrow=TRUE)</code> , then plot 1 will go in the upper left, 2 will go in the upper right, and 3 will go all the way across the bottom.

## Author(s)

David L. Borchers <[dlb@st-andrews.ac.uk](mailto:dlb@st-andrews.ac.uk)>

## Source

[http://www.cookbook-r.com/Graphs/Multiple\\_graphs\\_on\\_one\\_page\\_\(ggplot2\)/](http://www.cookbook-r.com/Graphs/Multiple_graphs_on_one_page_(ggplot2)/)

## Examples

```
if (require("ggplot2", quietly = TRUE)) {
  df <- data.frame(x = 1:10, y = cos(1:10), z = sin(1:10))
  p11 <- ggplot(data = df) +
    geom_line(mapping = aes(x, y), color = "red")
  p12 <- ggplot(data = df) +
    geom_line(mapping = aes(x, z), color = "blue")
  p13 <- ggplot(data = df) +
    geom_path(mapping = aes(y, z), color = "magenta")
  multiplot(
    p11, p12, p13,
    layout = rbind(c(1, 2), c(3, 3))
  )

  if (require("patchwork")) {
    (p11 + p12) / p13
  }
}
```

---

**plot.bru***Plot method for posterior marginals estimated by bru*

---

## Description

From version 2.11.0, `plot.bru(x, ...)` calls `plot.inla(x, ...)` from the INLA package, unless the first argument after `x` is a character, in which case the pre-2.11.0 behaviour is used, calling `plotmarginal.inla(x, ...)` instead.

Requires the `ggplot2` package.

## Usage

```
## S3 method for class 'bru'  
plot(x, ...)  
  
plotmarginal.inla(  
  result,  
  varname = NULL,  
  index = NULL,  
  link = function(x) {  
    x  
  },  
  add = FALSE,  
  ggp = TRUE,  
  lwd = 3,  
  ...  
)
```

## Arguments

<code>x</code>	a fitted <code>bru()</code> model.
<code>...</code>	Options passed on to other methods.
<code>result</code>	an <code>inla</code> or <code>bru</code> result object
<code>varname</code>	character; name of the variable to plot
<code>index</code>	integer; index of the random effect to plot
<code>link</code>	function; link function to apply to the variable
<code>add</code>	logical; if TRUE, add to an existing plot
<code>ggp</code>	logical; unused
<code>lwd</code>	numeric; line width

## Examples

```
## Not run:
if (require("ggplot2", quietly = TRUE)) {
  # Generate some data and fit a simple model
  input.df <- data.frame(x = cos(1:10))
  input.df <- within(
    input.df,
    y <- 5 + 2 * x + rnorm(length(x), mean = 0, sd = 0.1)
  )
  fit <- bru(y ~ x, family = "gaussian", data = input.df)
  summary(fit)

  # Plot the posterior density of the model's x-effect
  plot(fit, "x")
}

## End(Not run)
```

**plot.bru\_prediction** *Plot prediction using ggplot2*

## Description

Generates a base ggplot2 using `ggplot()` and adds a geom for input `x` using `gg`.  
Requires the `ggplot2` package.

## Usage

```
## S3 method for class 'bru_prediction'
plot(x, y = NULL, ...)

## S3 method for class 'prediction'
plot(x, y = NULL, ...)
```

## Arguments

- `x` a prediction object.
- `y` Ignored argument but required for S3 compatibility.
- `...` Arguments passed on to `gg.prediction()`.

## Value

an object of class `gg`

## Examples

```

if (bru_safe_inla() &&
    require(sn, quietly = TRUE) &&
    require(ggplot2, quietly = TRUE)) {
  # Generate some data

  input.df <- data.frame(x = cos(1:10))
  input.df <- within(input.df, y <- 5 + 2 * cos(1:10) + rnorm(10, mean = 0, sd = 0.1))

  # Fit a model with fixed effect 'x' and intercept 'Intercept'

  fit <- bru(y ~ x, family = "gaussian", data = input.df)

  # Predict posterior statistics of 'x'

  xpost <- predict(fit, NULL, formula = ~x_latent)

  # The statistics include mean, standard deviation, the 2.5% quantile, the median,
  # the 97.5% quantile, minimum and maximum sample drawn from the posterior as well as
  # the coefficient of variation and the variance.

  xpost

  # For a single variable like 'x' the default plotting method invoked by gg() will
  # show these statistics in a fashion similar to a box plot:
  ggplot() +
    gg(xpost)

  # The predict function can also be used to simultaneously estimate posteriors
  # of multiple variables:

  xipost <- predict(fit,
    newdata = NULL,
    formula = ~ c(
      Intercept = Intercept_latent,
      x = x_latent
    )
  )
  xipost

  # If we still want a plot in the previous style we have to set the bar parameter to TRUE

  p1 <- ggplot() +
    gg(xipost, bar = TRUE)
  p1

  # Note that gg also understands the posterior estimates generated while running INLA

  p2 <- ggplot() +
    gg(fit$summary.fixed, bar = TRUE)
  multiplot(p1, p2)

```

```

# By default, if the prediction has more than one row, gg will plot the column 'mean' against
# the row index. This is for instance useful for predicting and plotting function
# but not very meaningful given the above example:

ggplot() +
  gg(xipost)

# For ease of use we can also type

plot(xipost)

# This type of plot will show a ribbon around the mean, which visualizes the upper and lower
# quantiles mentioned above (2.5 and 97.5%). Plotting the ribbon can be turned off using the
# \code{ribbon} parameter

ggplot() +
  gg(xipost, ribbon = FALSE)

# Much like the other geoms produced by gg we can adjust the plot using ggplot2 style
# commands, for instance

ggplot() +
  gg(xipost) +
  gg(xipost, mapping = aes(y = median), ribbon = FALSE, color = "red")
}

```

**plotsample***Create a plot sample.***Description**

Creates a plot sample on a regular grid with a random start location.

**Usage**

```
plotsample(spdf, boundary, x.ppn = 0.25, y.ppn = 0.25, nx = 5, ny = 5)
```

**Arguments**

<code>spdf</code>	A <code>SpatialPointsDataFrame</code> defining the points that are to be sampled by the plot sample.
<code>boundary</code>	A <code>SpatialPolygonsDataFrame</code> defining the survey boundary within which the points occur.
<code>x.ppn</code>	The proportion of the x-axis that is to be included in the plots.
<code>y.ppn</code>	The proportion of the y-axis that is to be included in the plots.
<code>nx</code>	The number of plots in the x-dimension.
<code>ny</code>	The number of plots in the y-dimension.

**Value**

A list with three components:

`plots` A `SpatialPolygonsDataFrame` object containing the plots that were sampled.  
`dets` A `SpatialPointsDataFrame` object containing the locations of the points within the plots.  
`counts` A data frame containing the following columns

- `x` The x-coordinates of the centres of the plots within the boundary.
- `y` The y-coordinates of the centres of the plots within the boundary.
- `n` The numbers of points in each plot.
- `area` The areas of the plots within the boundary

**Examples**

```
# Some features require the raster package
if (bru_safe_sp() &&
    require("sp") &&
    require("raster", quietly = TRUE) &&
    require("ggplot2", quietly = TRUE) &&
    require("terra", quietly = TRUE) &&
    require("sf", quietly = TRUE)) {
  gorillas <- gorillas_sp()
  plotpts <- plotsample(gorillas$plots, gorillas$boundary,
    x.ppn = 0.4, y.ppn = 0.4, nx = 5, ny = 5
  )
  ggplot() +
    gg(plotpts$plots) +
    gg(plotpts$dets, pch = "+", cex = 2) +
    gg(gorillas$boundary)
}
```

point2count

*Convert a plot sample of points into one of counts.*

**Description**

Converts a plot sample with locations of each point within each plot, into a plot sample with only the count within each plot.

**Usage**

```
point2count(plots, dets)
```

**Arguments**

- plots            A SpatialPolygonsDataFrame object containing the plots that were sampled.  
 dets            A SpatialPointsDataFrame object containing the locations of the points within  
                 the plots.

**Value**

A SpatialPolygonsDataFrame with counts in each plot contained in slot @data\$n.

**Examples**

```
# Some features require the raster package
if (bru_safe_sp() &&
  require("sp") &&
  require("raster", quietly = TRUE) &&
  require("ggplot2", quietly = TRUE) &&
  require("terra", quietly = TRUE) &&
  require("sf", quietly = TRUE)) {
  gorillas <- gorillas_sp()
  plotpts <- plotsample(gorillas$plots, gorillas$boundary,
    x.ppn = 0.4, y.ppn = 0.4, nx = 5, ny = 5
  )
  p1 <- ggplot() +
    gg(plotpts$plots) +
    gg(plotpts$dets) +
    gg(gorillas$boundary)
  countdata <- point2count(plotpts$plots, plotpts$dets)
  x <- sp::coordinates(countdata)[, 1]
  y <- sp::coordinates(countdata)[, 2]
  count <- countdata@data$n
  p2 <- ggplot() +
    gg(gorillas$boundary) +
    gg(plotpts$plots) +
    geom_text(aes(label = count, x = x, y = y))
  multiplot(p1, p2, cols = 2)
}
```

**Description**

Point data and count data, together with intensity function and expected counts for a homogeneous 1-dimensional Poisson process example.

**Usage**

`data(Poisson1_1D)`

## Format

The data contain the following R objects:

`lambda1_1D` A function defining the intensity function of a nonhomogeneous Poisson process.

Note that this function is only defined on the interval (0,55).

`E_nc1` The expected counts of the gridded data.

`pts1` The locations of the observed points (a data frame with one column, named `x`).

`countdata1` A data frame with three columns, containing the count data:

- × The grid cell midpoint.

- count The number of detections in the cell.

- exposure The width of the cell.

## Examples

```
if (require("ggplot2", quietly = TRUE)) {
  data(Poisson1_1D)
  ggplot(countdata1) +
    geom_point(data = countdata1, aes(x = x, y = count), col = "blue") +
    ylim(0, max(countdata1$count)) +
    geom_point(data = pts1, aes(x = x), y = 0.2, shape = "|", cex = 4) +
    geom_point(
      data = countdata1, aes(x = x), y = 0, shape = "+",
      col = "blue", cex = 4
    ) +
    xlab(expression(bold(s))) +
    ylab("count")
}
```

## Description

Point data and count data, together with intensity function and expected counts for a unimodal nonhomogeneous 1-dimensional Poisson process example.

## Usage

```
data(Poisson2_1D)
```

## Format

The data contain the following R objects:

**lambda2\_1D:** A function defining the intensity function of a nonhomogeneous Poisson process.

Note that this function is only defined on the interval (0,55).

**cov2\_1D:** A function that gives what we will call a 'habitat suitability' covariate in 1D space.

**E\_nc2** The expected counts of the gridded data.

**pts2** The locations of the observed points (a data frame with one column, named **x**).

**countdata2** A data frame with three columns, containing the count data:

**x** The grid cell midpoint.

**count** The number of detections in the cell.

**exposure** The width of the cell.

## Examples

```
if (require("ggplot2", quietly = TRUE)) {
  data(Poisson2_1D)
  p1 <- ggplot(countdata2) +
    geom_point(data = countdata2, aes(x = x, y = count), col = "blue") +
    ylim(0, max(countdata2$count, E_nc2)) +
    geom_point(
      data = countdata2, aes(x = x), y = 0, shape = "+",
      col = "blue", cex = 4
    ) +
    geom_point(
      data = data.frame(x = countdata2$x, y = E_nc2), aes(x = x),
      y = E_nc2, shape = "_", cex = 5
    ) +
    xlab(expression(bold(s))) +
    ylab("count")
  ss <- seq(0, 55, length.out = 200)
  lambda <- lambda2_1D(ss)
  p2 <- ggplot() +
    geom_line(
      data = data.frame(x = ss, y = lambda),
      aes(x = x, y = y), col = "blue"
    ) +
    ylim(0, max(lambda)) +
    geom_point(data = pts2, aes(x = x), y = 0.2, shape = "|", cex = 4) +
    xlab(expression(bold(s))) +
    ylab(expression(lambda(bold(s)))))
  multiplot(p1, p2, cols = 1)
}
```

---

Poisson3\_1D*1-Dimensional NonHomogeneous Poisson example.*

---

**Description**

Point data and count data, together with intensity function and expected counts for a multimodal nonhomogeneous 1-dimensional Poisson process example. Counts are given for two different gridded data interval widths.

**Usage**

```
data(Poisson3_1D)
```

**Format**

The data contain the following R objects:

`lambda3_1D` A function defining the intensity function of a nonhomogeneous Poisson process.  
 Note that this function is only defined on the interval (0,55).  
`E_nc3a` The expected counts of gridded data for the wider bins (10 bins).  
`E_nc3b` The expected counts of gridded data for the wider bins (20 bins).  
`pts3` The locations of the observed points (a data frame with one column, named `x`).  
`countdata3a` A data frame with three columns, containing the count data for the 10-interval case:  
`countdata3b` A data frame with three columns, containing the count data for the 20-interval case:  
`x` The grid cell midpoint.  
`count` The number of detections in the cell.  
`exposure` The width of the cell.

**Examples**

```
if (require("ggplot2", quietly = TRUE)) {
  data(Poisson3_1D)
  # first the plots for the 10-bin case:
  p1a <- ggplot(countdata3a) +
    geom_point(data = countdata3a, aes(x = x, y = count), col = "blue") +
    ylim(0, max(countdata3a$count, E_nc3a)) +
    geom_point(
      data = countdata3a, aes(x = x), y = 0, shape = "+",
      col = "blue", cex = 4
    ) +
    geom_point(
      data = data.frame(x = countdata3a$x, y = E_nc3a),
      aes(x = x), y = E_nc3a, shape = "_", cex = 5
    ) +
    xlab(expression(bold(s))) +
    ylab("count")
  ss <- seq(0, 55, length.out = 200)
```

```

lambda <- lambda3_1D(ss)
p2a <- ggplot() +
  geom_line(
    data = data.frame(x = ss, y = lambda), aes(x = x, y = y),
    col = "blue"
  ) +
  ylim(0, max(lambda)) +
  geom_point(data = pts3, aes(x = x), y = 0.2, shape = "|", cex = 4) +
  xlab(expression(bold(s))) +
  ylab(expression(lambda(bold(s))))
multiplot(p1a, p2a, cols = 1)

# Then the plots for the 20-bin case:
p1a <- ggplot(countdata3b) +
  geom_point(data = countdata3b, aes(x = x, y = count), col = "blue") +
  ylim(0, max(countdata3b$count, E_nc3b)) +
  geom_point(
    data = countdata3b, aes(x = x), y = 0, shape = "+",
    col = "blue", cex = 4
  ) +
  geom_point(
    data = data.frame(x = countdata3b$x, y = E_nc3b),
    aes(x = x), y = E_nc3b, shape = "_", cex = 5
  ) +
  xlab(expression(bold(s))) +
  ylab("count")
ss <- seq(0, 55, length.out = 200)
lambda <- lambda3_1D(ss)
p2a <- ggplot() +
  geom_line(
    data = data.frame(x = ss, y = lambda), aes(x = x, y = y),
    col = "blue"
  ) +
  ylim(0, max(lambda)) +
  geom_point(data = pts3, aes(x = x), y = 0.2, shape = "|", cex = 4) +
  xlab(expression(bold(s))) +
  ylab(expression(lambda(bold(s))))
multiplot(p1a, p2a, cols = 1)
}

```

## Description

Takes a fitted `bru` object produced by the function `bru\(\)` and produces predictions given a new set of values for the model covariates or the original values used for the model fit. The predictions can be based on any R expression that is valid given these values/covariates and the joint posterior of the estimated random effects.

## Usage

```
## S3 method for class 'bru'
predict(
  object,
  newdata = NULL,
  formula = NULL,
  n.samples = 100,
  seed = 0L,
  probs = c(0.025, 0.5, 0.975),
  num.threads = NULL,
  used = NULL,
  drop = FALSE,
  ...,
  data = deprecated(),
  include = deprecated(),
  exclude = deprecated()
)
```

## Arguments

<code>object</code>	An object obtained by calling <code>bru()</code> or <code>lgcp()</code> .
<code>newdata</code>	A <code>data.frame</code> or <code>SpatialPointsDataFrame</code> of covariates needed for the prediction.
<code>formula</code>	A formula where the right hand side defines an R expression to evaluate for each generated sample. If <code>NULL</code> , the latent and hyperparameter states are returned as named list elements. See Details for more information.
<code>n.samples</code>	Integer setting the number of samples to draw in order to calculate the posterior statistics. The default is rather low but provides a quick approximate result.
<code>seed</code>	Random number generator seed passed on to <code>inla.posterior.sample</code>
<code>probs</code>	A numeric vector of probabilities with values in $[0, 1]$ , passed to <code>stats::quantile</code>
<code>num.threads</code>	Specification of desired number of threads for parallel computations. Default <code>NULL</code> , leaves it up to INLA. When <code>seed != 0</code> , overridden to "1:1"
<code>used</code>	Either <code>NULL</code> or a <code>bru_used()</code> object. Default <code>NULL</code> , uses auto-detection of used variables in the formula.
<code>drop</code>	logical; If <code>keep=FALSE</code> , <code>newdata</code> is a <code>Spatial*DataFrame</code> , and the prediction summary has the same number of rows as <code>newdata</code> , then the output is a <code>Spatial*DataFrame</code> object. Default <code>FALSE</code> .
<code>...</code>	Additional arguments passed on to <code>inla.posterior.sample()</code>
<code>data</code>	<b>[Deprecated]</b> Use <code>newdata</code> instead.
<code>include, exclude</code>	<b>[Deprecated]</b> If auto-detection of used variables fails, use <code>used</code> instead.

## Details

Mean value predictions are accompanied by the standard errors, upper and lower 2.5% quantiles, the median, variance, coefficient of variation as well as the variance and minimum and maximum sample value drawn in course of estimating the statistics.

Internally, this method calls [generate.bru\(\)](#) in order to draw samples from the model.

In addition to the component names (that give the effect of each component evaluated for the input data), the suffix \_latent variable name can be used to directly access the latent state for a component, and the suffix function \_eval can be used to evaluate a component at other input values than the expressions defined in the component definition itself, e.g. `field_eval(cbind(x, y))` for a component that was defined with `field(coordinates, ...)` (see also [bru\\_comp\\_eval\(\)](#)).

For "iid" models with `mapper = bm_index(n)`, `rnorm()` is used to generate new realisations for indices greater than `n`.

## Value

a `data.frame`, `sf`, or `Spatial*` object with predicted mean values and other summary statistics attached. Non-S4 object outputs have the class "bru\_prediction" added at the front of the class list.

## Examples

```
if (bru_safe_inla() &&
    require("sn", quietly = TRUE) &&
    require("ggplot2", quietly = TRUE) &&
    require("terra", quietly = TRUE) &&
    require("sf", quietly = TRUE)) {

  # Load the Gorilla data

  gorillas <- gorillas_sf

  # Plot the Gorilla nests, the mesh and the survey boundary

  ggplot() +
    gg(gorillas$mesh) +
    gg(gorillas$nests) +
    gg(gorillas$boundary, alpha = 0.1)

  # Define SPDE prior

  matern <- INLA::inla.spde2.pcmatern(
    gorillas$mesh,
    prior.sigma = c(0.1, 0.01),
    prior.range = c(0.01, 0.01)
  )

  # Define domain of the LCCP as well as the model components (spatial SPDE effect and Intercept)

  cmp <- geometry ~ field(geometry, model = matern) + Intercept(1)

  # Fit the model, with "eb" instead of full Bayes
```

```

fit <- lgcp(
  cmp,
  data = gorillas$nest,
  samplers = gorillas$boundary,
  domain = list(geometry = gorillas$mesh),
  options = list(control.inla = list(int.strategy = "eb")))
)

# Once we obtain a fitted model the predict function can serve various purposes.
# The most basic one is to determine posterior statistics of a univariate
# random variable in the model, e.g. the intercept

icpt <- predict(fit, NULL, ~ c(Intercept = Intercept_latent))
plot(icpt)

# The formula argument can take any expression that is valid within the model, for
# instance a non-linear transformation of a random variable

exp.icpt <- predict(fit, NULL, ~ c(
  "Intercept" = Intercept_latent,
  "exp(Intercept)" = exp(Intercept_latent)
))
plot(exp.icpt, bar = TRUE)

# The intercept is special in the sense that it does not depend on other variables
# or covariates. However, this is not true for the smooth spatial effects 'field'.
# In order to predict 'field' we have to define where (in space) to predict. For
# this purpose, the second argument of the predict function can take \code{data.frame}
# objects as well as Spatial objects. For instance, we might want to predict
# 'field' at the locations of the mesh vertices. Using

vrt <- fm_vertices(gorillas$mesh, format = "sf")

# we obtain these vertices as a SpatialPointsDataFrame

ggplot() +
  gg(gorillas$mesh) +
  gg(vrt, color = "red")

# Predicting 'mySmooth' at these locations works as follows

field <- predict(fit, vrt, ~field)

# Note that just like the input also the output will be a sf object with
# points and that the predicted statistics are simply added as columns

class(field)
head(vrt)
head(field)

# Plotting the mean, for instance, at the mesh node is straight forward

ggplot() +

```

```

gg(gorillas$mesh) +
  gg(field, aes(color = mean), size = 2)

# However, we are often interested in a spatial field and thus a linear interpolation,
# which can be achieved by using the gg mechanism for meshes

ggplot() +
  gg(gorillas$mesh, color = field$mean)

# Alternatively, we can predict the spatial field at a grid of locations, e.g. a
# sf object with a grid of points covering the relevant part of mesh

pxl <- fm_pixels(gorillas$mesh, format = "sf", mask = gorillas$boundary)
field2 <- predict(fit, p xl, ~field)

# This will give us a sf with the columns we are looking for

head(field2)
ggplot() +
  gg(gorillas$boundary) +
  gg(data = field2, geom = "tile")
}

```

**robins\_subset***robins\_subset*

## Description

This is the **robins\_subset** dataset, which is a subset of the full robins data set used to demonstrate a spatially varying trend coefficient model in Meehan et al. 2019. The dataset includes American Robin counts, along with time, location, and effort information, from Audubon Christmas Bird Counts (CBC) conducted in six US states between 1987 and 2016.

## Usage

```
robins_subset
```

## Format

The data are a data.frame with variables

**circle:** Four-letter code of the CBC circle.

**bcr:** Numeric code for the bird conservation region encompassing the count circle.

**state:** US state encompassing the count circle.

**year:** calendar year the count was conducted.

**std\_yr:** transformed year, with 2016 = 0.

**count:** number of robins recorded.

**log\_hrs:** the natural log of party hours.  
**lon:** longitude of the count circle centroid.  
**lat:** latitude of the count circle centroid.  
**obs:** unique record identifier.

## Source

<https://github.com/tmeeha/inlaSVCBC>

## References

Meehan, T.D., Michel, N.L., and Rue, H. 2019. Spatial modeling of Audubon Christmas Bird Counts reveals fine-scale patterns and drivers of relative abundance trends. *Ecosphere*, 10(4), p.e02707.

## Examples

```
if (require(ggplot2, quietly = TRUE)) {
  data(robins_subset, package = "inlabru") # get the data

  # plot the counts for one year of data
  ggplot(robins_subset[robins_subset$std_yr == 0, ]) +
    geom_point(aes(lon, lat, colour = count + 1)) +
    scale_colour_gradient(low = "blue", high = "red", trans = "log")
}
```

sample.lgcp

*Sample from an inhomogeneous Poisson process*

## Description

This function provides point samples from one- and two-dimensional inhomogeneous Poisson processes. The log intensity has to be provided via its values at the nodes of an `fm_mesh_1d` or `fm_mesh_2d` object. In between mesh nodes the log intensity is assumed to be linear.

## Usage

```
sample.lgcp(
  mesh,
  loglambda,
  strategy = NULL,
  R = NULL,
  samplers = NULL,
  ignore.CRS = FALSE
)
```

## Arguments

<code>mesh</code>	An <code>fmesher::fm_mesh_1d</code> or <code>fmesher::fm_mesh_2d</code> object
<code>loglambda</code>	vector or matrix; A vector of log intensities at the mesh vertices (for higher order basis functions, e.g. for <code>fm_mesh_1d</code> meshes, <code>loglambda</code> should be given as <code>mesh\$m</code> basis function weights rather than the values at the <code>mesh\$n</code> vertices) A single scalar is expanded to a vector of the appropriate length. If a matrix is supplied, one process sample for each column is produced.
<code>strategy</code>	Only relevant for 2D meshes. One of 'triangulated', 'rectangle', 'sliced-spherical', 'spherical'. The 'rectangle' method is only valid for CRS-less flat 2D meshes. If <code>NULL</code> or 'auto', the the likely fastest method is chosen; 'rectangle' for flat 2D meshes with no CRS, 'sliced-spherical' for CRS 'longlat' meshes, and 'triangulated' for all other meshes.
<code>R</code>	Numerical value only applicable to spherical and geographical meshes. It is interpreted as <code>R</code> is the equivalent Earth radius, in km, used to scale the lambda intensity. For CRS enabled meshes, the default is 6371. For CRS-less spherical meshes, the default is 1.
<code>samplers</code>	A <code>SpatialPolygonsDataFrame</code> or <code>fm_mesh_2d</code> object. Simulated points that fall outside these polygons are discarded.
<code>ignore.CRS</code>	logical; if <code>TRUE</code> , ignore any CRS information in the mesh. Default <code>FALSE</code> . This affects <code>R</code> and the permitted values for <code>strategy</code> .

## Details

For 2D processes on a sphere the `R` parameter can be used to adjust to sphere's radius implied by the mesh. If the intensity is very high the standard `strategy` "spherical" can cause memory issues. Using the "sliced-spherical" strategy can help in this case.

- For crs-less meshes on R2: Lambda is interpreted in the raw coordinate system. Output has an NA CRS.
- For crs-less meshes on S2: Lambda with raw units, after scaling the mesh to radius `R`, if specified. Output is given on the same domain as the mesh, with an NA CRS.
- For crs meshes on R2: Lambda is interpreted as per  $\text{km}^2$ , after scaling the globe to the Earth radius 6371 km, or `R`, if specified. Output given in the same CRS as the mesh.
- For crs meshes on S2: Lambda is interpreted as per  $\text{km}^2$ , after scaling the globe to the Earth radius 6371 km, or `R`, if specified. Output given in the same CRS as the mesh.

## Value

A `data.frame` (1D case), `SpatialPoints` (2D flat and 3D spherical surface cases) `SpatialPoints-DataFrame` (2D/3D surface cases with multiple samples). For multiple samples, the `data.frame` output has a column 'sample' giving the index for each sample. object of point locations.

## Author(s)

Daniel Simpson <[dp.simpson@gmail.com](mailto:dp.simpson@gmail.com)> (base rectangle and spherical algorithms), Fabian E. Bachl <[bach1fab@gmail.com](mailto:bach1fab@gmail.com)> (inclusion in `inlabru`, sliced spherical sampling), Finn Lindgren <[finn.lindgren@gmail.com](mailto:finn.lindgren@gmail.com)> (extended CRS support, triangulated sampling)

## Examples

```
# The INLA package is required
if (bru_safe_inla() &&
    bru_safe_sp() &&
    require("sp")) {
  vertices <- seq(0, 3, by = 0.1)
  mesh <- fm_mesh_1d(vertices)
  loglambda <- 5 - 0.5 * vertices
  pts <- sample.lgcp(mesh, loglambda)
  pts$y <- 0
  plot(vertices, exp(loglambda), type = "l", ylim = c(0, 150))
  points(pts, pch = "|")
}

# The INLA package is required
if (bru_safe_inla() &&
    require(ggplot2, quietly = TRUE) &&
    bru_safe_sp() &&
    require("sp") &&
    require("terra", quietly = TRUE) &&
    require("sf", quietly = TRUE)) {
  gorillas <- gorillas_sp()
  pts <- sample.lgcp(gorillas$mesh,
    loglambda = 1.5,
    samplers = gorillas$boundary
  )
  ggplot() +
    gg(gorillas$mesh) +
    gg(pts)
}
```

shrimp

*Blue and red shrimp in the Western Mediterranean Sea*

## Description

Blue and red shrimp in the Western Mediterranean Sea.

## Usage

```
data(shrimp)
```

## Format

A list of objects:

**hauls:** An sf object containing haul locations  
**mesh:** An fm\_mesh\_2d object containing a Delaunay triangulation mesh (a type of discretization of continuous space) covering the haul locations.  
**catch** Catch in Kg.  
**landing** Landing in Kg.  
**depth** Mean depth (in metres) of the fishery haul.

## Source

Pennino, Maria Grazia. Personal communication.

## References

Pennino, M. G., Paradinas, I., Munoz, F., Illian, J., Quilez-Lopez, A., Bellido, J.M., Conesa, D. Accounting for preferential sampling in species distribution models. *Ecology and Evolution*, 9(1), p653-663, 2019 [doi:10.1002/ece3.4789](https://doi.org/10.1002/ece3.4789)

## Examples

```
if (require(ggplot2, quietly = TRUE)) {
  data(shrimp, package = "inlabru", envir = environment())
  ggplot() +
    geom_fm(data = shrimp$mesh) +
    gg(shrimp$hauls, aes(col = catch)) +
    coord_sf(datum = fm_crs(shrimp$hauls))
}
```

**spde.posterior**

*Posteriors of SPDE hyper parameters and Matern correlation or covariance function.*

## Description

Calculate posterior distribution of the range, log(range), variance, or log(variance) parameter of a model's SPDE component. Can also plot Matern correlation or covariance function. `inla.spde.result`.

## Usage

```
spde.posterior(result, name, what = "range", quantile = 0.95)
```

## Arguments

<code>result</code>	An object inheriting from <code>inla</code> .
<code>name</code>	Character stating the name of the SPDE effect, see <code>names(result\$summary.random)</code> .
<code>what</code>	One of "range", "log.range", "variance", "log.variance", "matern.correlation" or "matern.covariance".
<code>quantile</code>	The target credible probability. Default 0.95.

**Value**

A prediction object.

**Author(s)**

Finn Lindgren <Finn.Lindgren@ed.ac.uk>

**Examples**

```
if (bru_safe_inla() && require(ggplot2, quietly = TRUE)) {  
  
  # Load 1D Poisson process data  
  
  data(Poisson2_1D, package = "inlabru")  
  
  # Take a look at the point (and frequency) data  
  
  ggplot(pts2) +  
    geom_histogram(aes(x = x), binwidth = 55 / 20, boundary = 0, fill = NA, color = "black") +  
    geom_point(aes(x), y = 0, pch = "|", cex = 4) +  
    coord_fixed(ratio = 1)  
  
  # Fit an LGCP model with  and SPDE component  
  
  x <- seq(0, 55, length.out = 20)  
  mesh1D <- fm_mesh_1d(x, boundary = "free")  
  mdl <- x ~ spde1D(x, model = INLA::inla.spde2.matern(mesh1D, constr = TRUE)) + Intercept(1)  
  fit <- lgcp(mdl, data = pts2, domain = list(x = mesh1D))  
  
  # Calculate and plot the posterior range  
  
  range <- spde.posterior(fit, "spde1D", "range")  
  plot(range)  
  
  # Calculate and plot the posterior log range  
  
  lrange <- spde.posterior(fit, "spde1D", "log.range")  
  plot(lrange)  
  
  # Calculate and plot the posterior variance  
  
  variance <- spde.posterior(fit, "spde1D", "variance")  
  plot(variance)  
  
  # Calculate and plot the posterior log variance  
  
  lvariance <- spde.posterior(fit, "spde1D", "log.variance")  
  plot(lvariance)  
  
  # Calculate and plot the posterior Matern correlation
```

```

matcor <- spde.posterior(fit, "spde1D", "matern.correlation")
plot(matcor)

# Calculate and plot the posterior Matern covariance

matcov <- spde.posterior(fit, "spde1D", "matern.covariance")
plot(matcov)
}

```

**summary.bru***Summary for an inlabru fit***Description**

Takes a fitted **bru** object produced by **bru()** or **lgcp()** and creates various summaries from it.

**Usage**

```

## S3 method for class 'bru'
summary(object, verbose = FALSE, ...)

## S3 method for class 'summary_bru'
print(x, ...)

```

**Arguments**

<b>object</b>	An object obtained from a <b>bru()</b> or <b>lgcp()</b> call
<b>verbose</b>	logical; If TRUE, include more details of the component definitions. If FALSE, only show basic component definition information. Default: FALSE
<b>...</b>	arguments passed on to component summary functions, see <b>summary.bru_comp()</b> .
<b>x</b>	An object to be printed

**Examples**

```

if (bru_safe_inla()) {
  # Simulate some covariates x and observations y
  input.df <- data.frame(x = cos(1:10))
  input.df <- within(input.df, {
    y <- 5 + 2 * x + rnorm(10, mean = 0, sd = 0.1)
  })
  
  # Fit a Gaussian likelihood model
  fit <- bru(y ~ x + Intercept(1), family = "gaussian", data = input.df)
  
  # Obtain summary
  fit$summary.fixed
}

```

```

if (bru_safe_inla()) {
  # Alternatively, we can use the bru_obs() function to construct the likelihood:

  lik <- bru_obs(family = "gaussian",
                  formula = y ~ x + Intercept,
                  data = input.df)
  fit <- bru(~ x + Intercept(1), lik)
  fit$summary.fixed
}

# An important addition to the INLA methodology is bru's ability to use
# non-linear predictors. Such a predictor can be formulated via bru_obs()'s
# \code{formula} parameter. The z(1) notation is needed to ensure that
# the z component should be interpreted as single latent variable and not
# a covariate:

if (bru_safe_inla()) {
  z <- 2
  input.df <- within(input.df, {
    y <- 5 + exp(z) * x + rnorm(10, mean = 0, sd = 0.1)
  })
  lik <- bru_obs(
    family = "gaussian", data = input.df,
    formula = y ~ exp(z) * x + Intercept
  )
  fit <- bru(~ z(1) + Intercept(1), lik)

  # Check the result (z posterior should be around 2)
  fit$summary.fixed
}

```

summary.bru\_obs

*Summary and print methods for observation models*

## Description

Summary and print methods for observation models

## Usage

```

## S3 method for class 'bru_obs'
summary(object, verbose = TRUE, ...)

## S3 method for class 'bru_obs_list'
summary(object, verbose = TRUE, ...)

## S3 method for class 'summary_bru_obs'

```

```

print(x, ...)

## S3 method for class 'summary_bru_obs_list'
print(x, ...)

## S3 method for class 'bru_obs'
print(x, ...)

## S3 method for class 'bru_obs_list'
print(x, ...)

```

### Arguments

<code>object</code>	Object to operate on
<code>verbose</code>	logical; If TRUE, include more details of the component definitions. If FALSE, only show basic component definition information. Default: TRUE
<code>...</code>	Arguments passed on to other summary methods
<code>x</code>	Object to be printed

### See Also

[bru\\_obs\(\)](#)

### Examples

```

obs <- bru_obs(y ~ ., data = data.frame(y = rnorm(10)))
summary(obs)
print(obs)

```

`summary.bru_options`    *Print inlabru options*

### Description

Print inlabru options

### Usage

```

## S3 method for class 'bru_options'
summary(
  object,
  legend = TRUE,
  include_global = TRUE,
  include_default = TRUE,
  ...
)

```

```
## S3 method for class 'summary_bru_options'
print(x, ...)
```

### Arguments

object	A <a href="#">bru_options</a> object to be summarised
legend	logical; If TRUE, include explanatory text, Default: TRUE
include_global	logical; If TRUE, include global override options
include_default	logical; If TRUE, include default options
...	Further parameters, currently ignored
x	A <a href="#">summary_bru_options</a> object to be printed

### Examples

```
if (interactive()) {
  options <- bru_options(verbose = TRUE)

  # Don't print options only set in default:
  print(options, include_default = FALSE)

  # Only include options set in the object:
  print(options, include_default = FALSE, include_global = FALSE)
}
```

toygroups

*Simulated 1D animal group locations and group sizes*

### Description

This data set serves to teach the concept of modelling species that gather in groups and where the grouping behaviour depends on space.

### Usage

```
data(toygroups)
```

### Format

The data are a list that contains these elements:

**groups:** A `data.frame` of group locations `x` and size `size`  
**df.size:** IGNORE THIS  
**df.intensity:** A `data.frame` with Poisson process intensity `d.lambda` at locations `x`  
**df.rate:** A `data.frame` the locations `x` and associated `rate` which parameterized the exponential distribution from which the group sizes were drawn.

## Examples

```
if (require(ggplot2, quietly = TRUE)) {
  # Load the data

  data("toygroups", package = "inlabru")

  # The data set is a simulation of animal groups residing in a 1D space. Their
  # locations in x-space are sampled from a Cox process with intensity

  ggplot(toygroups$df.intensity) +
    geom_line(aes(x = x, y = g.lambda))

  # Adding the simulated group locations to this plot we obtain

  ggplot(toygroups$df.intensity) +
    geom_line(aes(x = x, y = g.lambda)) +
    geom_point(data = toygroups$groups, aes(x, y = 0), pch = "|")

  # Each group has a size mark attached to it.
  # These group sizes are sampled from an exponential distribution
  # for which the rate parameter depends on the x-coordinate

  ggplot(toygroups$groups) +
    geom_point(aes(x = x, y = size))

  ggplot(toygroups$df.rate) +
    geom_line(aes(x, rate))
}
```

*toypoints*

*Simulated 2D point process data*

## Description

This data set serves as an example for basic inlabru.

## Usage

```
data(toypoints)
```

## Format

The data are a list that contains these elements:

- points An sf object of point locations and z measurements
- mesh An fm\_mesh\_2d object
- boundary An sf polygon denting the region of interest
- pred\_locs A sf object with prediction point locations

**Examples**

```
if (require("ggplot2")) {  
  ggplot() +  
    fmeshr::geom_fm(data = toypoints$mesh, alpha = 0) +  
    geom_sf(data = toypoints$boundary, fill = "blue", alpha = 0.1) +  
    geom_sf(data = toypoints$points, aes(color = z)) +  
    scale_color_viridis_c()  
}
```

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