

# Package ‘bayesnec’

April 21, 2022

**Title** A Bayesian No-Effect- Concentration (NEC) Algorithm

**Version** 2.0.2.4

**Description** Implementation of No-Effect-Concentration estimation that uses 'brms' (see Burkner (2017)<[doi:10.18637/jss.v080.i01](https://doi.org/10.18637/jss.v080.i01)>; Burkner (2018)<[doi:10.32614/RJ-2018-017](https://doi.org/10.32614/RJ-2018-017)>; Carpenter 'et al.' (2017)<[doi:10.18637/jss.v076.i01](https://doi.org/10.18637/jss.v076.i01)> to fit concentration(dose)-response data using Bayesian methods for the purpose of estimating 'ECX' values, but more particularly 'NEC' (see Fox (2010)<[doi:10.1016/j.ecoenv.2009.09.012](https://doi.org/10.1016/j.ecoenv.2009.09.012)>. This package expands and supersedes an original version implemented in R2jags, see Fisher, Ricardo and Fox (2020)<[doi:10.5281/ZENODO.3966864](https://doi.org/10.5281/ZENODO.3966864)>.

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**BugReports** <https://github.com/open-aims/bayesnec/issues>

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bayesnec-package      *The 'bayesnec' package.*

---

**Description**

A No-Effect-Concentration estimation package that uses brms (<https://github.com/paul-buerkner/brms>) to fit concentration (dose)-response data using Bayesian methods for the purpose of estimating both ECx values, but more particularly NEC. Please see ?bnec for more details.

**References**

Bürkner P-C (2018) Advanced Bayesian Multilevel Modeling with the R Package brms. The R Journal, 10: 395-411. doi:10.32614/RJ-2018-017.

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<code>+.bnecfit</code>	<i>"Add" multiple <code>bnecfit</code> objects into one single <code>bayesmanecfit</code> object containing Bayesian model averaging statistics.</i>
------------------------	--

---

### Description

"Add" multiple `bnecfit` objects into one single `bayesmanecfit` object containing Bayesian model averaging statistics.

### Usage

```
## S3 method for class 'bnecfit'
e1 + e2
```

### Arguments

`e1` An object of class `bnecfit`.  
`e2` An object of class `bnecfit`.

### Value

An object of class `bayesmanecfit`.

---

<code>amend</code>	<i>Amends an existing <code>bayesmanecfit</code> object, for example, by adding or removing fitted models.</i>
--------------------	--

---

### Description

Amends an existing `bayesmanecfit` object, for example, by adding or removing fitted models.

### Usage

```
amend(
  object,
  drop,
  add,
  loo_controls,
  x_range = NA,
  precision = 1000,
  sig_val = 0.01,
  priors
)
```

**Arguments**

object	An object of class <code>bayesmanecfit</code> , as returned by <code>bnec</code> .
drop	A <code>character</code> vector containing the names of model types you which to drop for the modified fit.
add	A <code>character</code> vector containing the names of model types to add to the modified fit.
loo_controls	A named <code>list</code> of two elements ("fitting" and/or "weights"), each being a named <code>list</code> containing the desired arguments to be passed on to <code>loo</code> (via "fitting") or to <code>loo_model_weights</code> (via "weights"). If "fitting" is provided with argument <code>pointwise = TRUE</code> (due to memory issues) and <code>family = "beta_binomial2"</code> , the <code>bnec</code> will fail because that is a custom family. If "weights" is not provided by the user, <code>bnec</code> will set the default method argument in <code>loo_model_weights</code> to "pseudobma". See <code>?loo_model_weights</code> for further info.
x_range	A range of predictor values over which to consider extracting ECx.
precision	The length of the predictor vector used for posterior predictions, and over which to extract ECx values. Large values will be slower but more precise.
sig_val	Probability value to use as the lower quantile to test significance of the predicted posterior values against the lowest observed concentration (assumed to be the control), to estimate NEC as an interpolated NOEC value from smooth ECx curves.
priors	An object of class <code>brmsprior</code> which specifies user-desired prior distributions of model parameters. If missing, <code>amend</code> will figure out a baseline prior for each parameter. It can also be specified as a named <code>list</code> where each name needs to correspond to the same string as model. See details.

**Value**

All successfully fitted `bayesmanecfit` model fits.

**Examples**

```
library(bayesnec)
data(manec_example)
exmp <- amend(manec_example, drop = "nec4param")
```

---

autoplot.bayesmanecfit

*autoplot.bayesmanecfit*

---

**Description**

`bayesnec` standard `ggplot2` plotting method.

**Usage**

```
## S3 method for class 'bayesianecfit'
autoplot(
  object,
  ...,
  nec = TRUE,
  ecx = FALSE,
  force_x = FALSE,
  xform = NA,
  all_models = FALSE,
  plot = TRUE,
  ask = TRUE,
  newpage = TRUE,
  multi_facet = TRUE
)
```

**Arguments**

object	An object of class <a href="#">bayesianecfit</a> as returned by function <a href="#">bnec</a> .
...	Additional arguments to be passed to <a href="#">ggbnec_data</a> .
nec	Should NEC values be added to the plot? Defaults to TRUE.
ecx	Should ECx values be added to the plot? Defaults to FALSE.
force_x	A <a href="#">logical</a> value indicating if the argument xform should be forced on the predictor values. This is useful when the user transforms the predictor beforehand (e.g. when using a non-standard base function).
xform	A function to apply to the returned estimated concentration values.
all_models	Should all individual models be plotted separately (defaults to FALSE) or should model averaged predictions be plotted instead?
plot	Should output <a href="#">ggplot</a> output be plotted? Only relevant if all = TRUE and multi_facet = FALSE.
ask	Indicates if the user is prompted before a new page is plotted. Only relevant if plot = TRUE and multi_facet = FALSE.
newpage	Indicates if the first set of plots should be plotted to a new page. Only relevant if plot = TRUE and multi_facet = FALSE.
multi_facet	Should all plots be plotted in one single panel via facets? Defaults to TRUE.

**Value**

A [ggplot](#) object.

**See Also**

Other autoplot methods: [autoplot.bayesianecfit\(\)](#)

## Examples

```
library(brms)
library(bayesnec)
options(mc.cores = 2)
data(nec_data)

necs <- bnec(y ~ crf(x, c("nec3param", "nec4param")), data = nec_data,
            iter = 2e2, family = Beta(link = "identity"))
nec3param <- pull_out(necs, "nec3param")
autoplot(nec3param)
autoplot(nec3param, nec = FALSE)
autoplot(nec3param, ecx = TRUE, ecx_val = 50)

# plot model averaged predictions
autoplot(necs)
# plot all panels together
autoplot(necs, ecx = TRUE, ecx_val = 50, all_models = TRUE)
# plots multiple models, one at a time, with interactive prompt
autoplot(necs, ecx = TRUE, ecx_val = 50, all_models = TRUE,
        multi_facet = FALSE)
```

---

autoplot.bayesnefit    *autoplot.bayesnefit*

---

## Description

[bayesnec](#) standard **ggplot2** plotting method.

## Usage

```
## S3 method for class 'bayesnefit'
autoplot(object, ..., nec = TRUE, ecx = FALSE, force_x = FALSE, xform = NA)
```

## Arguments

object	An object of class <a href="#">bayesnefit</a> as returned by function <a href="#">bnec</a> .
...	Additional arguments to be passed to <a href="#">ggbnec_data</a> .
nec	Should NEC values be added to the plot? Defaults to TRUE.
ecx	Should ECx values be added to the plot? Defaults to FALSE.
force_x	A <a href="#">logical</a> value indicating if the argument <code>xform</code> should be forced on the predictor values. This is useful when the user transforms the predictor beforehand (e.g. when using a non-standard base function).
xform	A function to apply to the returned estimated concentration values.

**Value**

A `ggplot` object.

**See Also**

Other autoplot methods: `autoplot.bayesianecfit()`

**Examples**

```
library(brms)
library(bayesnec)
options(mc.cores = 2)
data(nec_data)

necs <- bnecc(y ~ crf(x, c("nec3param", "nec4param")), data = nec_data,
             iter = 2e2, family = Beta(link = "identity"))
nec3param <- pull_out(necs, "nec3param")
autoplot(nec3param)
autoplot(nec3param, nec = FALSE)
autoplot(nec3param, ecx = TRUE, ecx_val = 50)

# plot model averaged predictions
autoplot(necs)
# plot all panels together
autoplot(necs, ecx = TRUE, ecx_val = 50, all_models = TRUE)
# plots multiple models, one at a time, with interactive prompt
autoplot(necs, ecx = TRUE, ecx_val = 50, all_models = TRUE,
         multi_facet = FALSE)
```

---

average\_endpoints      *average\_endpoints*

---

**Description**

Extracts posterior predicted endpoint values from a list of class `bayesnecfit` or `bayesianecfit` model fits and calculates a geometric mean.

**Usage**

```
average_endpoints(
  x,
  endpoint = "nec",
  ecx_val = 10,
  posterior = FALSE,
  type = "absolute",
  hormesis_def = "control",
  sig_val = 0.01,
```



```

precision = 1000,
x_range = NA,
xform = NA,
prob_vals = c(0.5, 0.025, 0.975)
)

```

## Arguments

x	A named <a href="#">list</a> of objects of class <a href="#">bayesnecfit</a> or <a href="#">bayesmanecfit</a> returned by <a href="#">bnec</a> .
endpoint	The type of endpoint to use in the mean. Takes values "nec", "ecx" or "nsec".
ecx_val	The desired percentage effect value. This must be a value between 1 and 99 (for type = "relative" and "absolute"), defaults to 10.
posterior	A <a href="#">logical</a> value indicating if the full posterior sample of calculated ECx values should be returned instead of just the median and 95 credible intervals.
type	A <a href="#">character</a> vector, taking values of "relative", "absolute" (the default) or "direct". See Details.
hormesis_def	A <a href="#">character</a> vector, taking values of "max" or "control". See Details.
sig_val	Probability value to use as the lower quantile to test significance of the predicted posterior values. against the lowest observed concentration (assumed to be the control), to estimate NEC as an interpolated NOEC value from smooth ECx curves.
precision	The number of unique x values over which to find ECx - large values will make the ECx estimate more precise.
x_range	A range of x values over which to consider extracting ECx.
xform	A function to apply to the returned estimated concentration values.
prob_vals	A vector indicating the probability values over which to return the estimated ECx value. Defaults to 0.5 (median) and 0.025 and 0.975 (95 percent credible intervals).

## Details

The geometric mean of values are simply the mean calculated on a log scale and back transformed through [exp](#), although we have added the capacity to accommodate zero values. Note that the function assumes that x has been modelled on the natural scale. Often C-R models are more stable on a log transformed or sqrt scaling. If the input [bayesnecfit](#) or [bayesmanecfit](#) model fits are already based on a re-scaling of the x (concentration) axis, it is important to pass an appropriate xform argument to ensure these are back transformed before the the geometric mean calculation is applied.

## Value

The geometric mean of the endpoints estimate values of the [bayesnecfit](#) or [bayesmanecfit](#) model fits contained in x. See Details.

**See Also**[bnec](#)**Examples**

```
## Not run:
library(brms)
library(bayesnec)
data(manec_example)
data(nec4param)
ecx4param <- pull_out(manec_example, model = "ecx4param")
average_endpoints(list("nec" = ecx4param, "ecx" = nec4param), ecx_val = 50)

## End(Not run)
```

---

bayesmanecfit-class    *Class bayesmanecfit of models fitted with the **brms** package*

---

**Description**

Multiple models fitted with the [bayesnec](#) package are represented as a bayesmanecfit object, which contains the original [brmsfit](#) fitted objects, names of non-linear models that were fitted, model averaging WAIC stats, sample size, mean posterior NEC values, mean model averaged predictions on the data scale, model averaged residuals, full posterior distribution of predicated values, and summary statistics of NEC statistics.

**Details**

See `methods(class = "bayesmanecfit")` for an overview of available methods.

**Slots**

`mod_fits` A [list](#) of fitted model outputs of class `prebayesnecfit` for each of the fitted models.  
`success_models` A [character](#) vector indicating the name of the successfully fitted models.  
`mod_stats` A [data.frame](#) of model fit statistics.  
`sample_size` The size of the posterior sample. information on the priors used in the model.  
`w_nec_posterior` The model-weighted posterior estimate of the NEC.  
`w_predicted_y` The model-weighted predicted values for the observed data.  
`w_residuals` Model-weighted residual values (i.e. observed - `w_predicted_y`).  
`w_pred_vals` A [list](#) containing model-weighted posterior predicted values based on the supplied precision and `x_range`.  
`w_nec` The summary stats (median and 95% credibility intervals) of `w_nec_posterior`.

**See Also**

[bayesnec](#), [bnec](#), [bayesnecfit](#)

---

bayesnefit-class	<i>Class bayesnefit of models fitted with the <b>brms</b> package</i>
------------------	---

---

### Description

Models fitted with the [bayesne](#) package are represented as a `bayesnefit` object, which contain the original `brmsfit` fitted object, list of initialisation values used, the validated `bayesneformula`, name of non-linear model that was fitted, posterior predictions, posterior parameter estimates and a series of other statistics.

### Details

See `methods(class = "bayesnefit")` for an overview of available methods.

### Slots

`fit` The fitted Bayesian model of class `brmsfit`.

`model` A `character` string indicating the name of the fitted model.

`inits` A `list` containing the initialisation values for to fit the model.

`bayesneformula` An object of class `bayesneformula` and `formula`.

`pred_vals` A `list` containing a `data.frame` of summary posterior predicted values and a vector containing based on the supplied precision and `x_range`.

`top` The estimate for parameter "top" in the fitted model.

`beta` The estimate for parameter "beta" in the fitted model.

`nec` The estimated NEC.

`f` The estimate for parameter "f" in the fitted model, NA if absent for the fitted model type.

`bot` The estimate for parameter "bot" in the fitted model, NA if absent for the fitted model type.

`d` The estimate for parameter "d" in the fitted model, NA if absent for the fitted model type.

`slope` The estimate for parameter "slope" in the fitted model, NA if absent for the fitted model type.

`ec50` The estimate for parameter "ec50" in the fitted model, NA if absent for the fitted model type.

`dispersion` An estimate of dispersion.

`predicted_y` The predicted values for the observed data.

`residuals` Residual values of the observed data from the fitted model.

`nec_posterior` A full posterior estimate of the NEC.

### See Also

[bayesne](#), [bnec](#), [bayesmanecfit](#), [bayesneformula](#)

---

bayesnecformula	<i>Set up a model formula for use in <a href="#">bayesnec</a></i>
-----------------	---

---

## Description

Set up a model formula for use in the [bayesnec](#) package allowing to define linear and non-linear (potentially multilevel) concentration-response models.

## Usage

```
bayesnecformula(formula, ...)
```

## Arguments

formula	Either a <a href="#">character</a> string defining an R formula or an actual <a href="#">formula</a> object. See details.
...	Unused.

## Details

See `methods(class = "bayesnecformula")` for an overview of available methods.

### General formula syntax

The formula argument accepts formulas of the following syntax:

```
response | aterms ~ crf(x,model) + glterms
```

### The population-level term: crf

[bayesnec](#) uses a special internal term called `crf` which sets the concentration-response equation to be evaluated based on some `x` predictor. The equation itself is defined by the argument `"model"`: a [character](#) vector containing a specific model, a concatenation of specific models, or a single string defining a particular group of models (or group of equations, see [models](#)). Internally this argument is substituted by an actual [brmsformula](#) which is then passed onto [brm](#) for model fitting.

### Group-level terms: glterms

The user has three options to define group-level effects in a [bayesnecformula](#): 1) a general "offset" group-level effect defined by the term `ogl` (as in e.g. `ogl(group_variable)`). This adds an additional population-level parameter `ogl` to the model defined by `crf`, analogously to what an intercept-only group-level effect would be in a classic linear model. 2) a group-level effect applied to all parameters in a model at once. This is done by the special term `pgl`, (as in e.g. `pgl(group_variable)`) which comes in handy so the user does not need to know the internal syntax and name of each parameter in the model. 3) a more classic approach where the user can specify which specific parameters — NB: that requires prior knowledge on the model structure and parameter names — to vary according to a grouping variable (as in e.g. `(bot | group_variable)`). [bayesnecformula](#) will ignore this term should the parameter not exist in the specified model or model suite. For example, the parameter `bot` exists in model `"nec4param"` but not in `"nec3param"`, so if the user specifies `model = "nec"` in `crf`, the term `(bot | group_variable)` will be dropped in models where that parameter does not exist.

**Further brms terms (largely untested)**

Currently `bayesnecformula` is quite agnostic about additional terms that are valid for a `brmsformula`. These are `aterms` and `ptersms` (see `?brmsformula`). The only capability that `bayesnecformula` does not allow is the addition of `ptersms` outside of the term `crf`. Although `ptersms` can be passed to predictor `x` within `crf`, we strongly discourage the user because those functionalities have not been tested yet. If this is extremely important to your research, please raise an issue on GitHub and we will consider. Currently, the only two `aterms` that have validated behaviour are:

1. `trials()` which is essential in binomially-distributed data, e.g. `y | trials(trials_variable)`, and 2) `weights`, e.g. `y | weights(weights_variable)`, following `brms` formula syntax. Please note that `brms` does not implement design weights as in other standard `base` function. From their help page, `brms` "takes the weights literally, which means that an observation with weight 2 receives 2 times more weight than an observation with weight 1. It also means that using a weight of 2 is equivalent to adding the corresponding observation twice to the data frame." Other `aterms` might be added, though we cannot attest to their functionality within `bayesnec`, i.e. checks will be done outside via `brm`.

**NB:** `aterms` other than `trials()` and `weights()` are currently omitted from `model.frame` output. If you need other `aterms` as part of that output please raise an issue on our GitHub page.

**Validation of formula** Please note that the function only checks for the input nature of the formula argument and adds a new class. This function **does not** perform any validation on the model nor checks on its adequacy to work with other functions in the package. For that please refer to the function `check_formula` which requires the dataset associated with the formula.

**Value**

An object of class `bayesnecformula` and `formula`.

**See Also**

`check_formula`, `model.frame`, `models`, `show_params`, `make_brmsformula`

**Examples**

```
library(bayesnec)

bayesnecformula(y ~ crf(x, "nec3param"))
# or use shot alias bnf
bayesnecformula(y ~ crf(x, "nec3param")) == bnf(y ~ crf(x, "nec3param"))
bnf(y | trials(tr) ~ crf(sqrt(x), "nec3param"))
bnf(y | trials(tr) ~ crf(x, "nec3param") + ogl(group_1) + pgl(group_2))
bnf(y | trials(tr) ~ crf(x, "nec3param") + (nec + top | group_1))

# complex transformations are not advisable because
# they are passed directly to Stan via brms
# and are likely to fail -- transform your variable beforehand!
try(bnf(y | trials(tr) ~ crf(scale(x), scale = TRUE), "nec3param")))
```

---

beta_binomial2	<i>Custom beta-binomial family</i>
----------------	------------------------------------

---

**Description**

Custom beta-binomial family

**Format**

An object of class `customfamily`

---

beta_binomial2_lpmf	<i>beta_binomial2_lpmf</i>
---------------------	----------------------------

---

**Description**

Beta-binomial wrapper LPMF

**Usage**

```
beta_binomial2_lpmf(y, mu, phi, trials)
```

**Arguments**

y	vector of observation successes.
mu	posterior mu.
phi	posterior phi.
trials	vector of observation trials.

**Value**

A `numeric` value or vector containing the probability density of the beta binomial distribution

---

beta_binomial2_rng	<i>beta_binomial2_rng</i>
--------------------	---------------------------

---

**Description**

Beta-binomial wrapper RNG

**Usage**

```
beta_binomial2_rng(mu, phi, trials)
```

**Arguments**

mu	posterior mu.
phi	posterior phi.
trials	vector of observation trials.

**Value**

A [numeric](#) value or vector containing random values of the beta binomial distribution

---

b nec	<i>b nec</i>
-------	--------------

---

**Description**

Fits a variety of NEC models using Bayesian analysis and provides a model averaged predictions based on WAIC model weights

**Usage**

```
b nec(
  formula,
  data,
  x_range = NA,
  precision = 1000,
  sig_val = 0.01,
  loo_controls,
  x_var = NULL,
  y_var = NULL,
  trials_var = NULL,
  model = NULL,
  random = NULL,
  random_vars = NULL,
  ...
)
```

## Arguments

formula	Either a <a href="#">character</a> string defining an R formula or an actual <a href="#">formula</a> object. See <a href="#">bayesnecformula</a> and <a href="#">check_formula</a> .
data	A <a href="#">data.frame</a> containing the data to use with the formula.
x_range	A range of predictor values over which to consider extracting ECx.
precision	The length of the predictor vector used for posterior predictions, and over which to extract ECx values. Large values will be slower but more precise.
sig_val	Probability value to use as the lower quantile to test significance of the predicted posterior values against the lowest observed concentration (assumed to be the control), to estimate NEC as an interpolated NOEC value from smooth ECx curves.
loo_controls	A named <a href="#">list</a> of two elements ("fitting" and/or "weights"), each being a named <a href="#">list</a> containing the desired arguments to be passed on to <a href="#">loo</a> (via "fitting") or to <a href="#">loo_model_weights</a> (via "weights"). If "fitting" is provided with argument <code>pointwise = TRUE</code> (due to memory issues) and <code>family = "beta_binomial2"</code> , the <a href="#">bnec</a> will fail because that is a custom family. If "weights" is not provided by the user, <a href="#">bnec</a> will set the default method argument in <a href="#">loo_model_weights</a> to "pseudobma". See <a href="#">?loo_model_weights</a> for further info.
x_var	Removed in version 2.0. Use formula instead. Used to be a <a href="#">character</a> indicating the column heading containing the predictor (concentration) variable.
y_var	Removed in version 2.0. Use formula instead. Used to be a <a href="#">character</a> indicating the column heading containing the response variable.
trials_var	Removed in version 2.0. Use formula instead. Used to be a <a href="#">character</a> indicating the column heading for the number of "trials" for binomial or beta_binomial2 response data, as it appears in "data" (if data is supplied).
model	Removed in version 2.0. Use formula instead. Used to be a <a href="#">character</a> vector indicating the model(s) to fit. See Details for more information.
random	Removed in version 2.0. Use formula instead. Used to be a named <a href="#">list</a> containing the random model formula to apply to model parameters.
random_vars	Removed in version 2.0. Use formula instead. Used to be a <a href="#">character</a> vector containing the names of the columns containing the variables used in the random model formula.
...	Further arguments to <a href="#">brm</a> .

## Details

### Overview

[bnec](#) serves as a wrapper for (currently) 23 (mostly) non-linear equations that are classically applied to concentration(dose)-response problems. The primary goal of these equations is to provide the user with estimates of No-Effect-Concentration (*NEC*), No-Significant-Effect-Concentration (*NSEC*), and Effect-Concentration (of specified percentage 'x', *ECx*) thresholds. These in turn are fitted through the [brm](#) function from package [brms](#) and therefore further arguments to [brm](#) are allowed. In the absence of those arguments, [bnec](#) makes its best attempt to calculate distribution family, priors and initialisation values for the user based on the characteristics of the data. Moreover, in the absence of user-specified values, [bnec](#) sets the number of iterations to 1e4 and warmup



period to  $\text{floor}(\text{iterations} / 5) * 4$ . The chosen models can be extended by the addition of **brms** special "aterms" as well as group-level effects. See [?bayesnecformula](#) for details.

### The available models/equations/formulas

The available equations (or models) can be found via the `models` function. Since version 2.0, `bnec` requires a specific formula structure which is fully explained in the help file of [bayesnecformula](#). This formula incorporates the information regarding the chosen model(s). If one single model is specified, `bnec` will return an object of class `bayesnecfit`; otherwise if model is either a concatenation of multiple models and/or a string indicating a family of models, `bnec` will return an object of class `bayesmanecfit`, providing they were successfully fitted. The major difference being that the output of the latter includes Bayesian model averaging statistics. These classes come with multiple associated methods such as `plot`, `autoplot`, `summary`, `print`, `model.frame` and `formula`.

model may also be one of "all", meaning all of the available models will be fit; "ecx" meaning only models excluding a specific NEC step parameter will be fit; "nec" meaning only models with a specific NEC step parameter will be fit; "bot\_free" meaning only models without a "bot" parameter (without a bottom plateau) will be fit; "zero\_bounded" are models that are bounded to be zero; or "decline" excludes all hormesis models, i.e., only allows a strict decline in response across the whole predictor range. Notice that if one of these group strings is provided together with a user-specified named list for the `brm`'s argument `prior`, the list names need to contain the actual model names, and not the group string, e.g. if `model = "ecx"` and `prior = my_priors` then `names(my_priors)` must contain `models("ecx")`. To check available models and associated parameters for each group, use the function `models` or to check the parameters of a specific model use the function `show_params`.

All models provide an estimate for NEC. For model types with "nec" as a prefix, NEC is directly estimated as parameter "nec" in the model. Models with "ecx" as a prefix are continuous curve models, typically used for extracting ECx values from concentration response data. In this instance the NEC value is defined as the concentration at which there is a user supplied (see argument `sig_val`) percentage certainty (based on the Bayesian posterior estimate) that the response falls below the estimated value of the upper asymptote (top) of the response (i.e. the response value is significantly lower than that expected in the case of no exposure). The default value for `sig_val` is 0.01, which corresponds to an alpha value of 0.01 for a one-sided test of significance.

### Further argument to `brm`

If not supplied via the `brm` argument `family`, the appropriate distribution will be guessed based on the characteristics of the input data. Guesses include: "bernoulli" / `bernoulli` / `bernoulli()`, "Beta" / `Beta` / `Beta()`, "binomial" / `binomial` / `binomial()`, "beta\_binomial2" / `beta_binomial2`, "Gamma" / `Gamma` / `Gamma()`, "gaussian" / `gaussian` / `gaussian()`, "negbinomial" / `negbinomial` / `negbinomial()`, or "poisson" / `poisson` / `poisson()`. Note, however, that "negbinomial" and "betabinomial2" require knowledge on whether the data is over-dispersed. As explained below in the Return section, the user can extract the dispersion parameter from a `bnec` call, and if they so wish, can refit the model using the "negbinomial" family.

Other families can be considered as required, please raise an [issue](#) on the GitHub development site if your required family is not currently available.

As a default, `bnec` sets the `brm` argument `sample_prior` to "yes" in order to sample draws from the priors in addition to the posterior distributions. Among others, these samples can be used to calculate Bayes factors for point hypotheses via [hypothesis](#).

Model averaging is achieved through a weighted sample of each fitted models posterior predictions, with weights derived using functions `loo` and `loo_model_weights` from **brms**. Argument to both

these functions can be passed via the `loo_controls` argument. Individual model fits can be pulled out for examination using function `pull_out`.

#### Additional technical notes

As some concentration-response data will use zero concentration which can cause numerical estimation issues, a small offset is added (1 / 10th of the next lowest value) to zero values of concentration where `x_var` are distributed on a continuous scale from 0 to infinity, or are bounded to 0, or 1.

#### NAs are thrown away

Stan's default behaviour is to fail when the input data contains NAs. For that reason **brms** excludes any NAs from input data prior to fitting, and does not allow them back in as is the case with e.g. `stats::lm` and `na.action = exclude`. So we advise that you exclude any NAs in your data prior to fitting because if you so wish that should facilitate merging predictions back onto your original dataset.

#### Value

If argument `model` is a single string, then an object of class `bayesnecfit`; if many strings or a set, an object of class `bayesmanecfit`.

#### See Also

[bayesnecformula](#), [check\\_formula](#), [models](#), [show\\_params](#)

#### Examples

```
## Not run:
library(bayesnec)
data(nec_data)

# A single model
exmp_a <- bnec(y ~ crf(x, model = "nec4param"), data = nec_data, chains = 2)
# Two models model
exmp_b <- bnec(y ~ crf(x, model = c("nec4param", "ecx4param")),
              data = nec_data, chains = 2)

## End(Not run)
```

---

bnecfit-class

*Class bnecfit of models fitted with function bnec*

---

#### Description

This is a wrapper class which will be attached to both `bayesnecfit` and `bayesmanecfit` classes. Useful for methods which might need to take either class as an input simultaneously.

#### Details

See `methods(class = "bnecfit")` for an overview of available methods.

**See Also**

[bayesnec](#), [bnec](#), [bayesnecfit](#), [bayesmanecfit](#)

---

c.bnecfif	<i>Concatenate multiple <a href="#">bnecfif</a> objects into one single <a href="#">bayesmanecfit</a> object containing Bayesian model averaging statistics.</i>
-----------	--

---

**Description**

Concatenate multiple [bnecfif](#) objects into one single [bayesmanecfit](#) object containing Bayesian model averaging statistics.

**Usage**

```
## S3 method for class 'bnecfif'
c(x, ...)
```

**Arguments**

x                   An object of class [bnecfif](#).  
 ...                 Additional objects of class [bnecfif](#).

**Value**

An object of class [bayesmanecfit](#).

---

check_chains	<i>check_chains</i>
--------------	---------------------

---

**Description**

Plots HMC chains for a [bayesnecfit](#) or [bayesmanecfit](#) model fit as returned by [bnec](#).

**Usage**

```
check_chains(x, ...)
```

**Arguments**

x                   An object of class [bayesnecfit](#) or [bayesmanecfit](#) as returned by [bnec](#).  
 ...                 arguments used when class is [bayesmanecfit](#).

**Value**

No return value, generates a plot or writes a pdf to file.

## Examples

```
library(bayesnec)
data(manec_example)

# print to device
check_chains(manec_example)
```

---

```
check_chains.bayesmanecfit
      check_chains.bayesmanecfit
```

---

## Description

Plots HMC chains for a [bayesnecfit](#) model fit as returned by [bnec](#).

## Usage

```
## S3 method for class 'bayesmanecfit'
check_chains(x, ..., filename = NA)
```

## Arguments

x	An object of class <a href="#">bayesnecfit</a> or <a href="#">bayesmanecfit</a> as returned by <a href="#">bnec</a> .
...	arguments used when class is <a href="#">bayesmanecfit</a> .
filename	An optional <a href="#">character</a> vector to be used as a pdf filename in the case of a <a href="#">bayesmanecfit</a> . Any non empty character string will indicate the user wants to save the plots.

## Value

No return value, generates a plot or writes a pdf to file.

## Examples

```
library(bayesnec)
data(manec_example)

# print to device
check_chains(manec_example)
```

---

```
check_chains.bayesnefit  
  check_chains.bayesnefit
```

---

**Description**

Plots HMC chains for a `bayesnefit` model fit as returned by `bnec`.

**Usage**

```
## S3 method for class 'bayesnefit'  
check_chains(x, ...)
```

**Arguments**

`x` An object of class `bayesnefit` or `bayesmanefit` as returned by `bnec`.  
`...` arguments used when class is `bayesmanefit`.

**Value**

No return value, generates a plot or writes a pdf to file.

**Examples**

```
library(bayesnefit)  
data(manec_example)  
  
# print to device  
check_chains(manec_example)
```

---

```
check_chains.default  check_chains.default
```

---

**Description**

Plots HMC chains for a `bayesnefit` or `bayesmanefit` model fit as returned by `bnec`.

**Usage**

```
## Default S3 method:  
check_chains(x, ...)
```

**Arguments**

`x` An object of class `bayesnefit` or `bayesmanefit` as returned by `bnec`.  
`...` arguments used when class is `bayesmanefit`.

**Value**

No return value, generates a plot or writes a pdf to file.

**Examples**

```
library(bayesnec)
data(manec_example)

# print to device
check_chains(manec_example)
```

---

check_formula	<i>Check if input model formula is appropriate to use with <a href="#">bayesnec</a></i>
---------------	---

---

**Description**

Perform a series of checks to ensure that the input formula is valid for usage within [bayesnec](#).

**Usage**

```
check_formula(formula, data, run_par_checks = FALSE)
```

**Arguments**

formula	An object of class <a href="#">bayesnecformula</a> as returned by function <a href="#">bayesnecformula</a> .
data	A <a href="#">data.frame</a> containing the variables specified in formula.
run_par_checks	See details. A <a href="#">logical</a> defining whether random terms for specific parameters should be checked against the underlying concentration-response model defined in formula. Defaults to FALSE.

**Details**

This function allows the user to make sure that the input formula will allow for a successful model fit with the function [bnec](#). Should all checks pass, the function returns the original formula. Otherwise it will fail and requires that the user fixes it until they're able to use it with [bnec](#).

The argument `run_par_checks` is irrelevant for most usages of this package because it only applies if three conditions are met: 1) the user has specified a group-level effect; 2) the group-level effects is parameter specific (e.g. `(par | group_variable)` rather than `pgl/ogl(group_variable)`); and 3) The user is keen to learn if the specified parameter is found in the specified model (via argument `model` in the `crf` term – see details in `?bayesnecformula`).

**NB:** `aterms` other than `trials()` and `weights()` are currently omitted from `model.frame` output. If you need other `aterms` as part of that output please raise an issue on our GitHub page. See details about `aterms` in `?bayesnecformula`.

**Value**

A validated object of class `bayesnecformula` and `formula`.

**See Also**

`bnec`, `bayesnecformula`

**Examples**

```
library(bayesnec)
nec3param <- function(beta, nec, top, x) {
  top * exp(-exp(beta) * (x - nec) *
    ifelse(x - nec < 0, 0, 1))
}

data <- data.frame(x = seq(1, 20, length.out = 10), tr = 100, wght = c(1, 2),
  group_1 = sample(c("a", "b"), 10, replace = TRUE),
  group_2 = sample(c("c", "d"), 10, replace = TRUE))
data$y <- nec3param(beta = -0.2, nec = 4, top = 100, data$x)

# returns error
# > f_1 <- y ~ crf(x, "nec3param") + z
# regular formula not allowed, wrap it with function bnf
# > check_formula(f_1, data)
# population-level covariates are not allowed
# > check_formula(bnf(f_1), data)

# expect a series of messages for because not all
# nec models have the "bot" parameter
f_2 <- y | trials(tr) ~ crf(x, "nec") + (nec + bot | group_1)
check_formula(bnf(f_2), data, run_par_checks = TRUE)

# runs fine
f_3 <- "y | trials(tr) ~ crf(sqrt(x), \"nec3param\")"
check_formula(bnf(f_3), data)
f_4 <- y | trials(tr) ~ crf(x, "nec3param") + ogl(group_1) + pgl(group_2)
inherits(check_formula(bnf(f_4), data), "bayesnecformula")
```

---

check\_priors

*Plots the prior and posterior parameter probability densities from an object of class `bayesnecfit` or `bayesmanecfit`.*

---

**Description**

Plots the prior and posterior parameter probability densities from an object of class `bayesnecfit` or `bayesmanecfit`.

**Usage**

```
check_priors(object, filename = NA)
```

**Arguments**

**object** An object of class `bayesnecfit` or `bayesmanecfit` returned by `b nec`.

**filename** An optional `character` vector to be used as a pdf filename in the case of a `bayesmanecfit`. Any non empty character string will indicate the user wants to save the plots.

**Value**

A plot of the prior and posterior parameter probability densities.

**See Also**

`b nec`

**Examples**

```
library(bayesnec)
data(manec_example)
check_priors(manec_example)
```

---

`compare_endpoints`      *compare\_endpoints*

---

**Description**

Extracts posterior predicted values from a list of class `bayesnecfit` or `bayesmanecfit` model fits and compares these via bootstrap re sampling.

**Usage**

```
compare_endpoints(
  x,
  comparison = "nec",
  ecx_val = 10,
  type = "absolute",
  hormesis_def = "control",
  sig_val = 0.01,
  precision,
  x_range = NA
)
```



**Arguments**

x	A named <a href="#">list</a> of objects of class <a href="#">bayesnecfit</a> or <a href="#">bayesmanecfit</a> returned by <a href="#">bnec</a> .
comparison	The posterior predictions to compare, takes values of "nec", "nsec", "ecx" or "fitted".
ecx_val	The desired percentage effect value. This must be a value between 1 and 99 (for type = "relative" and "absolute"), defaults to 10.
type	A <a href="#">character</a> vector, taking values of "relative", "absolute" (the default) or "direct". See <a href="#">Details</a> .
hormesis_def	A <a href="#">character</a> vector, taking values of "max" or "control". See <a href="#">Details</a> .
sig_val	Probability value to use as the lower quantile to test significance of the predicted posterior values. against the lowest observed concentration (assumed to be the control), to estimate NEC as an interpolated NOEC value from smooth ECx curves.
precision	The number of unique x values over which to find ECx - large values will make the ECx estimate more precise.
x_range	A range of x values over which to consider extracting ECx.

**Value**

A named [list](#) containing bootstrapped differences in posterior predictions of the [bayesnecfit](#) or [bayesmanecfit](#) model fits contained in x. See [Details](#).

**See Also**

[bnec](#)

**Examples**

```
## Not run:
library(bayesnec)
data(manec_example)
nec4param <- pull_out(manec_example, model = "nec4param")
ecx4param <- pull_out(manec_example, model = "ecx4param")
compare_endpoints(list("nec" = ecx4param, "ecx" = nec4param), ecx_val = 50)

## End(Not run)
```

---

compare_fitted	<i>compare_fitted</i>
----------------	-----------------------

---

### Description

Extracts posterior predicted values from a list of class `bayesnecfit` or `bayesmanecfit` model fits and compares these across a vector of fitted values.

### Usage

```
compare_fitted(x, precision = 50, x_range = NA)
```

### Arguments

<code>x</code>	A named <code>list</code> of objects of class <code>bayesnecfit</code> or <code>bayesmanecfit</code> returned by <code>bne</code> .
<code>precision</code>	The number of unique <code>x</code> values over which to find ECx - large values will make the ECx estimate more precise.
<code>x_range</code>	A range of <code>x</code> values over which to consider extracting ECx.

### Value

A named `list` containing bootstrapped differences in posterior predictions of the `bayesnecfit` or `bayesmanecfit` model fits contained in `x`. See Details.

### See Also

`bne`

### Examples

```
## Not run:  
library(bayesnec)  
data(manec_example)  
nec4param <- pull_out(manec_example, model = "nec4param")  
ecx4param <- pull_out(manec_example, model = "ecx4param")  
compare_fitted(list("nec" = ecx4param, "ecx" = nec4param))  
  
## End(Not run)
```

---

compare\_posterior      *compare\_posterior*

---

## Description

Extracts posterior predicted values from a list of class `bayesnecfit` or `bayesmanecfit` model fits and compares these via bootstrap re sampling.

## Usage

```
compare_posterior(
  x,
  comparison = "nec",
  ecx_val = 10,
  type = "absolute",
  hormesis_def = "control",
  sig_val = 0.01,
  precision,
  x_range = NA
)
```

## Arguments

<code>x</code>	A named <a href="#">list</a> of objects of class <code>bayesnecfit</code> or <code>bayesmanecfit</code> returned by <code>bneec</code> .
<code>comparison</code>	The posterior predictions to compare, takes values of "nec", "nsec", "ecx" or "fitted".
<code>ecx_val</code>	The desired percentage effect value. This must be a value between 1 and 99 (for type = "relative" and "absolute"), defaults to 10.
<code>type</code>	A <a href="#">character</a> vector, taking values of "relative", "absolute" (the default) or "direct". See Details.
<code>hormesis_def</code>	A <a href="#">character</a> vector, taking values of "max" or "control". See Details.
<code>sig_val</code>	Probability value to use as the lower quantile to test significance of the predicted posterior values. against the lowest observed concentration (assumed to be the control), to estimate NEC as an interpolated NOEC value from smooth ECx curves.
<code>precision</code>	The number of unique x values over which to find ECx - large values will make the ECx estimate more precise.
<code>x_range</code>	A range of x values over which to consider extracting ECx.

## Value

A named [list](#) containing bootstrapped differences in posterior predictions of the `bayesnecfit` or `bayesmanecfit` model fits contained in `x`. See Details.

**See Also**[bnec](#)**Examples**

```
## Not run:
library(bayesnec)
data(manec_example)
nec4param <- pull_out(manec_example, model = "nec4param")
ecx4param <- pull_out(manec_example, model = "ecx4param")
compare_posterior(list("nec" = ecx4param, "ecx" = nec4param), ecx_val = 50)

## End(Not run)
```

---

**dispersion***Posterior dispersion*

---

**Description**

Calculates posterior dispersion metric

**Usage**

```
dispersion(model, summary = FALSE, seed = 10)
```

**Arguments**

model	An object of class <a href="#">bayesnecfit</a> whose distribution family is either <a href="#">poisson</a> or <a href="#">binomial</a> .
summary	Logical. Should summary stats be returned instead of full vector? Defaults to FALSE.
seed	Change seed for reproducible purposes.

**Details**

This function calculates a dispersion metric which takes the ratio between the observed relative to simulated Pearson residuals sums of squares.

**Value**

A [numeric](#) vector. If `summary` is FALSE, an n-long vector containing the dispersion metric, where n is the number of post warm-up posterior draws from the [brmsfit](#) object. If TRUE, then a [data.frame](#) containing the summary stats (mean, median, 95% highest density intervals) of the dispersion metric.

## References

Zuur, A. F., Hilbe, J. M., & Ieno, E. N. (2013). A Beginner's Guide to GLM and GLMM with R: A Frequentist and Bayesian Perspective for Ecologists. Highland Statistics Limited.

## Examples

```
library(bayesnec)
data(nec_data)
nec_data$y <- as.integer(round(nec_data$y * 100))
nec4param <- bnec(y ~ crf(x, "nec4param"), data = nec_data, chains = 2)
dispersion(nec4param, summary = TRUE)
```

---

ecx	<i>Extracts the predicted ECx value as desired from an object of class <a href="#">bayesnecfit</a> or <a href="#">bayesnecfit</a>.</i>
-----	--

---

## Description

Extracts the predicted ECx value as desired from an object of class [bayesnecfit](#) or [bayesnecfit](#).

## Usage

```
ecx(
  object,
  ecx_val = 10,
  precision = 1000,
  posterior = FALSE,
  type = "absolute",
  hormesis_def = "control",
  x_range = NA,
  xform = NA,
  prob_vals = c(0.5, 0.025, 0.975)
)
```

## Arguments

object	An object of class <a href="#">bayesnecfit</a> or <a href="#">bayesmanecfit</a> returned by <a href="#">bnec</a> .
ecx_val	The desired percentage effect value. This must be a value between 1 and 99 (for type = "relative" and "absolute"), defaults to 10.
precision	The number of unique x values over which to find ECx - large values will make the ECx estimate more precise.
posterior	A <a href="#">logical</a> value indicating if the full posterior sample of calculated ECx values should be returned instead of just the median and 95 credible intervals.
type	A <a href="#">character</a> vector, taking values of "relative", "absolute" (the default) or "direct". See Details.

hormesis_def	A <a href="#">character</a> vector, taking values of "max" or "control". See Details.
x_range	A range of x values over which to consider extracting ECx.
xform	A function to apply to the returned estimated concentration values.
prob_vals	A vector indicating the probability values over which to return the estimated ECx value. Defaults to 0.5 (median) and 0.025 and 0.975 (95 percent credible intervals).

### Details

type "relative" is calculated as the percentage decrease from the maximum predicted value of the response (top) to the minimum predicted value of the response. Type "absolute" (the default) is calculated as the percentage decrease from the maximum value of the response (top) to 0 (or bot for a 4 parameter model fit). Type "direct" provides a direct estimate of the x value for a given y. Note that for the current version, ECx for an "nechorme" (NEC Hormesis) model is estimated at a percent decline from the control. For `hormesis_def`, if "max", then ECx values are calculated as a decline from the maximum estimates (i.e. the peak at NEC); if "control", then ECx values are calculated relative to the control, which is assumed to be the lowest observed concentration.

### Value

A vector containing the estimated ECx value, including upper and lower 95% credible interval bounds.

### See Also

[bnec](#)

### Examples

```
library(brms)
library(bayesnec)
data(manec_example)
ecx(manec_example, ecx_val = 50)
ecx(manec_example)
```

---

expand\_manec

*Extracts a range of statistics from a list of [prebayesnecfit](#) objects.*

---

### Description

Extracts a range of statistics from a list of [prebayesnecfit](#) objects.

**Usage**

```

expand_manec(
  object,
  formula,
  x_range = NA,
  precision = 1000,
  sig_val = 0.01,
  loo_controls
)

```

**Arguments**

object	A <a href="#">list</a> of objects of class <a href="#">prebayesnecfit</a> .
formula	Either a <a href="#">character</a> string defining an R formula or an actual <a href="#">formula</a> object. See <a href="#">bayesnecformula</a> and <a href="#">check_formula</a> . It could also be a <a href="#">list</a> of formulas if multiple objects are passed to object.
x_range	A range of predictor values over which to consider extracting ECx.
precision	The length of the predictor vector used for posterior predictions, and over which to extract ECx values. Large values will be slower but more precise.
sig_val	Probability value to use as the lower quantile to test significance of the predicted posterior values against the lowest observed concentration (assumed to be the control), to estimate NEC as an interpolated NOEC value from smooth ECx curves.
loo_controls	A named <a href="#">list</a> of two elements ("fitting" and/or "weights"), each being a named <a href="#">list</a> containing the desired arguments to be passed on to <a href="#">loo</a> (via "fitting") or to <a href="#">loo_model_weights</a> (via "weights"). If "fitting" is provided with argument <code>pointwise = TRUE</code> (due to memory issues) and <code>family = "beta_binomial2"</code> , the <a href="#">bnec</a> will fail because that is a custom family. If "weights" is not provided by the user, <a href="#">bnec</a> will set the default method argument in <a href="#">loo_model_weights</a> to "pseudobma". See <a href="#">?loo_model_weights</a> for further info.

**Value**

A [list](#) of model statistical output derived from the input model list.

---

expand\_nec

*Extracts a range of statistics from a [prebayesnecfit](#) object.*

---

**Description**

Extracts a range of statistics from a [prebayesnecfit](#) object.

**Usage**

```

expand_nec(
  object,
  formula,
  x_range = NA,
  precision = 1000,
  sig_val = 0.01,
  loo_controls,
  ...
)

```

**Arguments**

object	An object of class <code>prebayesnecfit</code> .
formula	Either a <code>character</code> string defining an R formula or an actual <code>formula</code> object. See <code>bayesnecformula</code> and <code>check_formula</code> .
x_range	A range of predictor values over which to consider extracting ECx.
precision	The length of the predictor vector used for posterior predictions, and over which to extract ECx values. Large values will be slower but more precise.
sig_val	Probability value to use as the lower quantile to test significance of the predicted posterior values against the lowest observed concentration (assumed to be the control), to estimate NEC as an interpolated NOEC value from smooth ECx curves.
loo_controls	A named <code>list</code> of two elements ("fitting" and/or "weights"), each being a named <code>list</code> containing the desired arguments to be passed on to <code>loo</code> (via "fitting") or to <code>loo_model_weights</code> (via "weights"). If "fitting" is provided with argument <code>pointwise = TRUE</code> (due to memory issues) and <code>family = "beta_binomial2"</code> , the <code>bnec</code> will fail because that is a custom family. If "weights" is not provided by the user, <code>bnec</code> will set the default method argument in <code>loo_model_weights</code> to "pseudobma". See <code>?loo_model_weights</code> for further info.
...	Further arguments to internal function.

**Value**

A `list` of model statistical output derived from the input model object.

---

formula.bayesmanecfit *formula.bayesmanecfit*

---

**Description**

formula.bayesmanecfit



**Usage**

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

**Arguments**

x                    An object of class `bayesmanecfit` as returned by `bneec`.

...                  Further arguments passed to or from other methods.

model                A `character` string indicating which model or suite of models to pull out.

**Value**

An object of class `formula`.

---

formula.bayesnecfit    *formula.bayesnecfit*

---

**Description**

formula.bayesnecfit

**Usage**

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

**Arguments**

x                    An object of class `bayesnecfit` as returned by `bneec`.

...                  Further arguments passed to or from other methods.

**Value**

An object of class `formula`.

---

ggbnec\_data                      *Creates the data.frame for plotting with [autoplot](#).*

---

## Description

Creates the data.frame for plotting with [autoplot](#).

## Usage

```
ggbnec_data(  
  x,  
  add_nec = TRUE,  
  add_ecx = FALSE,  
  force_x = FALSE,  
  xform = NA,  
  ...  
)
```

## Arguments

x	An object of class <a href="#">bayesnecfit</a> or <a href="#">bayesmanecfit</a> , as returned by function <a href="#">bnc</a> .
add_nec	Should NEC values be added to the plot? Defaults to TRUE.
add_ecx	Should ECx values be added to the plot? Defaults to FALSE.
force_x	A <a href="#">logical</a> value indicating if the argument xform should be forced on the predictor values. This is useful when the user transforms the predictor beforehand (e.g. when using a non-standard base function).
xform	A function to apply to the returned estimated concentration values.
...	Additional arguments to be passed to <a href="#">ecx</a> . By default, function <a href="#">ecx</a> returns EC10.

## Value

A [data.frame](#).

## Examples

```
library(bayesnec)  
options(mc.cores = 2)  
data(manec_example)  
  
ggbnec_data(manec_example)  
ggbnec_data(manec_example, add_ecx = TRUE, ecx_val = 50)
```

---

herbicide

*Herbicide phytotoxicity data*

---

## Description

Herbicide phytotoxicity dataset from Jones & Kerswell (2003).

## Format

An object of class `data.frame` with 580 rows and 3 columns.

## Details

The response data (Fv/Fm) Chlorophyll fluorescence measurements of symbiotic dinoflagellates still in the host tissue of the coral (in hospite or in vivo) were measured using a DIVING-PAM chlorophyll fluorometer (Walz) on vertical planes of tissue 2 to 3 cm above the base of the corals, using either a 6 mm (*Acropora formosa*) or 2 mm (*Seriatopora hystrix*) fibre-optic probe. Parameters measured were the maximum potential quantum yield (Fv/Fm).

Additional information on each of the herbicides included is available from the original publication Jones & Kerswell (2003).

The columns are as follows:

**herbicide** The herbicide (chr).

**concentration** The treatment concentration in  $\mu\text{g} / \text{L}$  (dbl).

**fvfm** Maximum effective quantum yield (dbl).

## References

Jones RJ, Kerswell AP (2003) Phytotoxicity of Photosystem II (PSII) herbicides to coral. *Marine Ecology Progress Series*, 261: 149-159. doi: 10.3354/meps261149.

## Examples

```
head(herbicide)
```

---

is_manecsummary	<i>Checks if argument is a manecsummary object</i>
-----------------	--

---

**Description**

Checks if argument is a manecsummary object

**Usage**

```
is_manecsummary(x)
```

**Arguments**

x                    An R object

**Value**

A [logical](#)

---

is_necsummary	<i>Checks if argument is a necsummary object</i>
---------------	--

---

**Description**

Checks if argument is a necsummary object

**Usage**

```
is_necsummary(x)
```

**Arguments**

x                    An R object

---

```
log_lik_beta_binomial2
      log_lik_beta_binomial2
```

---

**Description**

Beta-binomial wrapper LL

**Usage**

```
log_lik_beta_binomial2(i, prep)
```

**Arguments**

i	observation i.
prep	data with posterior.

**Value**

Log likelihood of the beta binomial distribution

---

```
make_brmsformula      Expose the final brmsformula
```

---

**Description**

Checks the input formula according to [bayesnec](#) requirements and expose the final [brmsformula](#) which is to be fitted via package **brms**.

**Usage**

```
make_brmsformula(formula, data)
```

**Arguments**

formula	Either a <a href="#">character</a> string defining an R formula or an actual <a href="#">formula</a> object. See details.
data	A <a href="#">data.frame</a> containing the variables specified in formula.

**Value**

A named [list](#), with each element containing the final [brmsformula](#) to be passed to [brm](#).

**See Also**

[bayesnecformula](#), [check\\_formula](#)

## Examples

```

library(bayesnec)
nec3param <- function(beta, nec, top, x) {
  top * exp(-exp(beta) * (x - nec) *
    ifelse(x - nec < 0, 0, 1))
}

data <- data.frame(x = seq(1, 20, length.out = 10), tr = 100, wght = c(1, 2),
  group_1 = sample(c("a", "b"), 10, replace = TRUE),
  group_2 = sample(c("c", "d"), 10, replace = TRUE))
data$y <- nec3param(beta = -0.2, nec = 4, top = 100, data$x)

# make one single model
f_1 <- "y | trials(tr) ~ crf(sqrt(x), \"nec3param\")"
make_brmsformula(f_1, data)
# make an entire class of models
f_2 <- y ~ crf(x, "ecx") + ogl(group_1) + pgl(group_2)
make_brmsformula(f_2, data)

```

---

manecsummary-class      *Class manecsummary of models fitted with the **brms** package*

---

## Description

Multiple models fitted with the [bayesnec](#) package are summarised as a manecsummary object, which contains the name of the non-linear models fitted, the family distribution used to fit all the models, the total post-warm-up sample size, a table containing the model weights, the method to calculate the weights, whether this model is an ECx-type model (see details below), and the ECx summary values should the user decide to calculate them.

## Details

See `methods(class = "manecsummary")` for an overview of available methods.

## Slots

`models` A [character](#) string indicating the name of the fitted non-linear models.

`family` A [list](#) indicating the family distribution and link function used to fit all the models.

`sample_size` The total post-warm-up sample size.

`mod_weights` A table containing the model weights.

`mod_weights_method` The method to calculate the weights.

`ecx_mods` A [logical](#) indicating which models are ECx-type models.

`nec_vals` The model-averaged NEC values. Note that if model stack contains ECx-type models, these will be via NSEC proxies.

`ecs` A [list](#) containing the ECx values should the user decide to calculate them (see the non-exported `bayesnec::summary.bayesnecfit` help file for details). Different from the single-model case of class `bayesnecfit`, these ECx estimates will be based on the model weights.

`rhat_issues` A [list](#) detailing whether each fitted model exhibited convergence issues based on the Rhat evaluation.

### See Also

[bayesnec](#), [bnec](#), [bayesnecfit](#), [bayesmanecfit](#), [necsummary](#)

---

manec_example	<i>Example bayesmanecfit object</i>
---------------	-------------------------------------

---

### Description

Example bayesmanecfit object

### Format

An object of class `bayesmanecfit`. This was created to reduce run time in examples and tests, and to give the user an example to toy with. This was fitted to `bayesnec` built-in mock dataset (see [?nec\\_data](#)), using models "nec4param" and "ecx4param". The number of chains were set to 2 and number of iterations were 50 only to make sure that package size was below 5 Mb. See help files for function `bnec` and class `bayesmanecfit` for details.

### Source

Code used to generate these models can be downloaded from [https://github.com/open-AIMS/bayesnec/blob/master/data-raw/manec\\_example.R](https://github.com/open-AIMS/bayesnec/blob/master/data-raw/manec_example.R)

---

model.frame.bayesmanecfit	<i>model.frame.bayesmanecfit</i>
---------------------------	----------------------------------

---

### Description

model.frame.bayesmanecfit

### Usage

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

**Arguments**

formula	An object of class <code>bayesmanecfit</code> as returned by <code>b nec</code> .
...	Further arguments passed to or from other methods.
model	A <code>character</code> string indicating which model or suite of models to pull out.

**Value**

A `data.frame` containing the data used to fit the model chosen from the existing `bayesmanecfit` set.

---

```
model.frame.bayesnecfit
      model.frame.bayesnecfit
```

---

**Description**

`model.frame.bayesnecfit`

**Usage**

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

**Arguments**

formula	An object of class <code>bayesnecfit</code> as returned by <code>b nec</code> .
...	Further arguments passed to or from other methods.

**Value**

A `data.frame` containing the data used to fit the model.

---

```
model.frame.bayesnecformula
      Extracting the model frame from a bayesnecformula
```

---

**Description**

Recovers evaluated `data.frame` given input data and a formula of class `bayesnecformula`.

**Usage**

```
## S3 method for class 'bayesnecformula'
model.frame(formula, data, ...)
```



**Arguments**

formula	A formula of class <code>bayesnecformula</code> .
data	A <code>data.frame</code> containing the variables specified in formula.
...	Additional arguments to be passed to <code>check_formula</code> .

**Details**

If the formula contains transformations to variables `x` and `y`, these are evaluated and returned as part of the `data.frame`.

**Value**

A `data.frame` with additional attributes detailing the population-level variables (attribute `"bnec_pop"`) (response `y`, predictor `x`, and, if binomial a formula, trials) and, if applicable, the group-level variables (attribute `"bnec_group"`).

**Examples**

```
library(bayesnec)
nec3param <- function(beta, nec, top, x) {
  top * exp(-exp(beta) * (x - nec) *
    ifelse(x - nec < 0, 0, 1))
}

data <- data.frame(x = seq(1, 20, length.out = 10), tr = 100, wght = c(1, 2),
  group_1 = sample(c("a", "b"), 10, replace = TRUE),
  group_2 = sample(c("c", "d"), 10, replace = TRUE))
data$y <- nec3param(beta = -0.2, nec = 4, top = 100, data$x)

f_1 <- y ~ crf(x, "nec3param")
f_2 <- "y | trials(tr) ~ crf(sqrt(x), \"nec3param\")"
f_3 <- y | trials(tr) ~ crf(x, "nec3param") + ogl(group_1) + pgl(group_2)
f_4 <- y | trials(tr) ~ crf(x, "nec3param") + (nec + top | group_1)

m_1 <- model.frame(bnf(f_1), data)
attr(m_1, "bnec_pop")
model.frame(bnf(f_2), data)
m_3 <- model.frame(bnf(f_3), data)
attr(m_3, "bnec_group")
model.frame(bnf(f_4), data)
```

---

models

*models*


---

**Description**

Lists the fitted or available models

**Usage**

```
models(object)
```

**Arguments**

**object** An object of class `bayesnecfit` or `bayesmanecfit` as returned by `bnec`, a `character` vector indicating the type of model set for which to list the available models, or a `numeric` vector indicating the natural range of values which the models should be able to handle (see details). If missing all available models and their groups are listed.

**Details**

The available models are "nec3param", "nec4param", "nechorme", "nechorme4", "necsigm", "neclin", "neclinhorme", "nechormepwr", "nechorme4pwr", "nechormepwr01", "ecxlin", "ecxexp", "ecxsigm", "ecx4param", "ecxwb1", "ecxwb2", "ecxwb1p3", "ecxwb2p3", "ecxll5", "ecxll4", "ecxll3", "ecxhorme4", and "ecxhorme5".

To see the model formula and parameters for a specific model use the function `show_params`.

To see all the models in an available set (e.g. "all", "nec" or "ecx") use the function `models` specifying the group name.

To see the model names, model formula and parameters fitted in an existing `bayesnecfit` or `bayesmanecfit` model object use the function `models` specifying the fitted object.

To see what models are available for a given type of data use the function `models` passing a `numeric` vector indicating the range of possible data types. Models that have an exponential decay (most models with parameter "beta") with no "bot" parameter are zero-bounded and are not suitable for the Gaussian family, or any family modelled using a logit or log link function. Models with a linear decay (containing the string "lin" in their name) are not suitable for modelling families that are zero bounded (Gamma, Poisson, Negative binomial) using an identity link. Models with a linear decay or hormesis linear increase (all models with parameter "slope") are not suitable for modelling families that are 0, 1 bounded (binomial, beta, betabinomial2). These restrictions do not need to be controlled by the user and a call to `bnec` with `models = "all"` will simply exclude inappropriate models.

**Value**

A `list` of the available or fitted models.

**Examples**

```
library(bayesnec)
# default to all models and model groups
models()
# single model
show_params("nec3param")
# group of models
models("all")
# models that are suitable for 0,1 bounded data
models(c(0,1))
```

---

nec	<i>Extracts the predicted NEC value as desired from an object of class <a href="#">bayesnecfit</a> or <a href="#">bayesmanecfit</a>.</i>
-----	--

---

### Description

Extracts the predicted NEC value as desired from an object of class [bayesnecfit](#) or [bayesmanecfit](#).

### Usage

```
nec(object, posterior = FALSE, xform = NA, prob_vals = c(0.5, 0.025, 0.975))
```

### Arguments

object	An object of class <a href="#">bayesnecfit</a> or <a href="#">bayesmanecfit</a> returned by <a href="#">bnec</a> .
posterior	A <a href="#">logical</a> value indicating if the full posterior sample of calculated NEC values should be returned instead of just the median and 95% credible intervals.
xform	A function to apply to the returned estimated concentration values.
prob_vals	A vector indicating the probability values over which to return the estimated NEC value. Defaults to 0.5 (median) and 0.025 and 0.975 (95 percent credible intervals).

### Value

A vector containing the estimated NEC value, including upper and lower 95% credible interval bounds (or other interval as specified by `prob_vals`).

### See Also

[bnec](#)

### Examples

```
library(bayesnec)
data(manec_example)
nec(manec_example)
```

---

necsummary-class	<i>Class necsummary of models fitted with the <b>brms</b> package</i>
------------------	---

---

### Description

Single models fitted with the [bayesnec](#) package are summarised as a `necsummary` object, which contains the original `brmsfit` object summary, the name of the non-linear model fitted, whether this model is an ECx-type model (see details below), and the ECx summary values should the user decide to calculate them.

### Details

See `methods(class = "necsummary")` for an overview of available methods.

### Slots

`brmssummary` The standard summary for the fitted Bayesian model of class `brmsfit`.

`model` A [character](#) string indicating the name of the fitted non-linear model.

`is_ecx` A [logical](#) indicating whether `model` is an ECx-type model.

`ecs` A [list](#) containing the ECx values should the user decide to calculate them (see the non-exported `bayesnec:::summary.bayesnecfit` help file for details).

### See Also

[bayesnec](#), [bnec](#), [bayesnecfit](#), [bayesmanecfit](#), [manecsummary](#)

---

nec_data	<i>Example data of non-linear decay</i>
----------	---

---

### Description

A simulated dataset containing a series of response measurements as a function of a concentration axis. Data simulated by Diego Barneche.

### Format

A data frame with 100 rows and 2 variables:

- x: Concentration (predictor) axis.
- y: Response.

---

nsec	<i>Extracts the predicted NSEC value as desired from an object of class <a href="#">bayesnecfit</a> or <a href="#">bayesmanecfit</a>.</i>
------	---

---

## Description

Extracts the predicted NSEC value as desired from an object of class [bayesnecfit](#) or [bayesmanecfit](#).

## Usage

```
nsec(
  object,
  sig_val = 0.01,
  precision = 1000,
  posterior = FALSE,
  x_range = NA,
  hormesis_def = "control",
  xform = NA,
  prob_vals = c(0.5, 0.025, 0.975)
)
```

## Arguments

object	An object of class <a href="#">bayesnecfit</a> or <a href="#">bayesmanecfit</a> returned by <a href="#">bnec</a> .
sig_val	Probability value to use as the lower quantile to test significance of the predicted posterior values. against the lowest observed concentration (assumed to be the control), to estimate NEC as an interpolated NOEC value from smooth ECx curves.
precision	The number of unique x values over which to find NSEC - large values will make the NSEC estimate more precise.
posterior	A <a href="#">logical</a> value indicating if the full posterior sample of calculated NSEC values should be returned instead of just the median and 95 credible intervals.
x_range	A range of x values over which to consider extracting NSEC.
hormesis_def	A <a href="#">character</a> vector, taking values of "max" or "control". See Details.
xform	A function to apply to the returned estimated concentration values.
prob_vals	A vector indicating the probability values over which to return the estimated NSEC value. Defaults to 0.5 (median) and 0.025 and 0.975 (95 percent credible intervals).

## Details

For `hormesis_def`, if "max", then NSEC values are calculated as a decline from the maximum estimates (i.e. the peak at nec); if "control", then ECx values are calculated relative to the control, which is assumed to be the lowest observed concentration.

**Value**

A vector containing the estimated NSEC value, including upper and lower 95% credible interval bounds.

**See Also**

[bnec](#)

**Examples**

```
library(bayesnec)

data(manec_example)
nsec(manec_example)
```

---

`plot.bayesmanecfit`     *plot.bayesmanecfit*

---

**Description**

Generates a plot of a fitted [bayesmanecfit](#) object, as returned by [bnec](#).

**Usage**

```
## S3 method for class 'bayesmanecfit'
plot(
  x,
  ...,
  CI = TRUE,
  add_nec = TRUE,
  position_legend = "topright",
  add_ec10 = FALSE,
  xform = NA,
  lxform = NA,
  force_x = FALSE,
  jitter_x = FALSE,
  jitter_y = FALSE,
  ylab = "Response",
  xlab = "Predictor",
  xticks = NA,
  all_models = FALSE
)
```

**Arguments**

x	An object of class <code>bayesnefit</code> as returned by <code>bnec</code> .
...	Additional arguments to <code>plot</code> .
CI	A <code>logical</code> value indicating if credibility intervals on the model fit should be plotted, calculated as the upper and lower bounds of the individual predicted values from all posterior samples.
add_nec	A <code>logical</code> value indicating if the estimated NEC value and 95% credible intervals should be added to the plot.
position_legend	A <code>numeric</code> vector indicating the location of the NEC or EC10 legend, as per a call to <code>legend</code> .
add_ec10	A <code>logical</code> value indicating if an estimated EC10 value and 95% credible intervals should be added to the plot.
xform	A function to be applied as a transformation of the x data.
lxform	A function to be applied as a transformation only to axis labels and the annotated NEC / EC10 values.
force_x	A <code>logical</code> value indicating if the argument <code>xform</code> should be forced on the predictor values. This is useful when the user transforms the predictor beforehand (e.g. when using a non-standard base function).
jitter_x	A <code>logical</code> value indicating if the x data points on the plot should be jittered.
jitter_y	A <code>logical</code> value indicating if the y data points on the plot should be jittered.
ylab	A <code>character</code> vector to use for the y-axis label.
xlab	A <code>character</code> vector to use for the x-axis label.
xticks	A <code>numeric</code> vector indicate where to place the tick marks of the x-axis.
all_models	A <code>logical</code> value indicating if all models in the model set should be plotted simultaneously, or if a model average plot should be returned.

**Value**

a plot of the fitted model

---

plot.bayesnefit      *plot.bayesnefit*

---

**Description**

Generates a plot of a fitted `bayesnefit` model, as returned by `bnec`.

**Usage**

```
## S3 method for class 'bayesnefit'
plot(
  x,
  ...,
  CI = TRUE,
  add_nec = TRUE,
  position_legend = "topright",
  add_ec10 = FALSE,
  xform = NA,
  lxform = NA,
  force_x = FALSE,
  jitter_x = FALSE,
  jitter_y = FALSE,
  ylab = "Response",
  xlab = "Predictor",
  xticks = NA
)
```

**Arguments**

x	An object of class <code>bayesnefit</code> as returned by <code>bnef</code> .
...	Additional arguments to <code>plot</code> .
CI	A <b>logical</b> value indicating if credibility intervals on the model fit should be plotted, calculated as the upper and lower bounds of the individual predicted values from all posterior samples.
add_nec	A <b>logical</b> value indicating if the estimated NEC value and 95% credible intervals should be added to the plot.
position_legend	A <b>numeric</b> vector indicating the location of the NEC or EC10 legend, as per a call to <code>legend</code> .
add_ec10	A <b>logical</b> value indicating if an estimated EC10 value and 95% credible intervals should be added to the plot.
xform	A function to be applied as a transformation of the x data.
lxform	A function to be applied as a transformation only to axis labels and the annotated NEC / EC10 values.
force_x	A <b>logical</b> value indicating if the argument <code>xform</code> should be forced on the predictor values. This is useful when the user transforms the predictor beforehand (e.g. when using a non-standard base function).
jitter_x	A <b>logical</b> value indicating if the x data points on the plot should be jittered.
jitter_y	A <b>logical</b> value indicating if the y data points on the plot should be jittered.
ylab	A <b>character</b> vector to use for the y-axis label.
xlab	A <b>character</b> vector to use for the x-axis label.
xticks	A numeric vector indicate where to place the tick marks of the x-axis.



**Value**

a plot of the fitted model

---

posterior\_epred\_beta\_binomial2  
*posterior\_epred\_beta\_binomial2*

---

**Description**

Beta-binomial wrapper posterior\_epred method

**Usage**

posterior\_epred\_beta\_binomial2(prepare)

**Arguments**

prepare            data with posterior.

**Value**

A numeric value or vector containing predicted random values of the beta binomial distribution

---

posterior\_predict\_beta\_binomial2  
*posterior\_predict\_beta\_binomial2*

---

**Description**

Beta-binomial wrapper posterior\_predict method

**Usage**

posterior\_predict\_beta\_binomial2(i, prepare, ...)

**Arguments**

i                    observation i.  
prepare            data with posterior.  
...                   unused.

**Value**

A numeric value or vector containing predicted probability values of the beta binomial distribution

---

```
prebayesianecfit-class Class prebayesianecfit of models fitted with the brms package
```

---

### Description

This is an intermediate class that was created to make both `bayesianecfit` and `bayesianecfit` objects lighter to handle. It contains the original `brmsfit` fitted object, name of non-linear model that was fitted, the list of initialisation values applied, and the validated `bayesianecformula`.

### Details

See `methods(class = "prebayesianecfit")` for an overview of available methods.

### Slots

`fit` The fitted Bayesian model of class `brmsfit`.  
`model` A `character` string indicating the name of the fitted model.  
`inits` A `list` containing the initialisation values for to fit the model.  
`bayesianecformula` An object of class `bayesianecformula` and `formula`.

### See Also

`bayesianec`, `bnec`, `bayesianecfit`, `bayesianecfit`, `bayesianecformula`

---

```
predict.bayesianecfit predict.bayesianecfit
```

---

### Description

`predict.bayesianecfit`

### Usage

```
## S3 method for class 'bayesianecfit'
predict(object, ..., precision = 100, x_range = NA)
```

### Arguments

<code>object</code>	An object of class <code>bayesianecfit</code> as returned by <code>bnec</code> .
<code>...</code>	Unused.
<code>precision</code>	A <code>numeric</code> vector of length 1 indicating the number of x values over which to predict values.
<code>x_range</code>	A <code>numeric</code> vector of length 2 indicating the range of x values over which to make predictions.

**Value**

A [list](#) containing two elements: a [data.frame](#) with predictor x and fitted y values plus lower and upper credible intervals; a [matrix](#) of M x N, with M being the number of posterior draws and N being the number of observations in the input data.

---

predict.bayesnefit    *predict.bayesnefit*

---

**Description**

predict.bayesnefit

**Usage**

```
## S3 method for class 'bayesnefit'
predict(object, ..., precision = 100, x_range = NA)
```

**Arguments**

object	An object of class <a href="#">bayesnefit</a> as returned by <a href="#">bnec</a> .
...	Unused.
precision	A <a href="#">numeric</a> vector of length 1 indicating the number of x values over which to predict values.
x_range	A <a href="#">numeric</a> vector of length 2 indicating the range of x values over which to make predictions.

**Value**

A [list](#) containing two elements: a [data.frame](#) with predictor x and fitted y values plus lower and upper credible intervals; a [matrix](#) of M x N, with M being the number of posterior draws and N being the number of observations in the input data.

---

print.bayesmanecfit    *print.bayesmanecfit*

---

**Description**

print.bayesmanecfit

**Usage**

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

**Arguments**

x                    An object of class `bayesmanecfit` as returned by `bneec`.  
 ...                    Further arguments to function `summary`.

**Value**

A `list` containing a summary of the model fit as returned by a `brmsfit` object for each model.

---

`print.bayesneecfit`      *print.bayesneecfit*

---

**Description**

`print.bayesneecfit`

**Usage**

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

**Arguments**

x                    An object of class `bayesneecfit` as returned by `bneec`.  
 ...                    Further arguments to function `summary`.

**Value**

A `list` containing a summary of the model fit as returned for a `brmsfit` object.

---

`print.manecsummary`      *print.manecsummary*

---

**Description**

`print.manecsummary`

**Usage**

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

**Arguments**

x                    An object of class `manecsummary` as returned by `summary.bayesmanecfit`.  
 ...                    Unused.

**Value**

A list containing a summary of model features and statistics.

---

```
print.necsummary      print.necsummary
```

---

**Description**

```
print.necsummary
```

**Usage**

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

**Arguments**

x	An object of class <code>necsummary</code> as returned by <code>summary.bayesnecfit</code> .
...	Unused.

**Value**

A `list` containing a summary of model features and statistics.

---

```
pull_out      pull_out
```

---

**Description**

Subsets model(s) from an existing object of class `bayesmanecfit`

**Usage**

```
pull_out(manec, model, loo_controls, ...)
```

**Arguments**

manec	An object of class <code>bayesmanecfit</code> as returned by <code>bneec</code> .
model	A <code>character</code> string indicating which model or suite of models to pull out.
loo_controls	A named <code>list</code> of two elements ("fitting" and/or "weights"), each being a named <code>list</code> containing the desired arguments to be passed on to <code>loo</code> (via "fitting") or to <code>loo_model_weights</code> (via "weights"). If "fitting" is provided with argument <code>pointwise = TRUE</code> (due to memory issues) and <code>family = "beta_binomial2"</code> , the <code>bneec</code> will fail because that is a custom family. If "weights" is not provided by the user, <code>bneec</code> will set the default method argument in <code>loo_model_weights</code> to "pseudobma". See <code>?loo_model_weights</code> for further info.
...	Additional arguments to <code>expand_nec</code> or <code>expand_manec</code> .

**Value**

If `model` is a string representing a single model, an object of class `bayesnecfit`; If `model` is instead a string depicting a suite of models, and object of class `bayesmanecfit`.

**See Also**

[bnec](#), [models](#).

**Examples**

```
## Not run:
library(bayesnec)
data(manec_example)
nec4param <- pull_out(manec_example, model = "nec4param")
# use "ecx" to get all ECx-containing models
# (only one ["ecx4param"] in this minimal example)
ecx_models <- pull_out(manec_example, model = "ecx")

## End(Not run)
```

---

`pull_prior`

*pull\_prior*

---

**Description**

Extracts the priors from an object of class `bayesnecfit` or `bayesmanecfit`.

**Usage**

```
pull_prior(object)
```

**Arguments**

`object` An object of class `bayesnecfit` or `bayesmanecfit` returned by `bnec`.

**Value**

A `list` containing the priors.

**Examples**

```
library(bayesnec)
data(manec_example)
pull_prior(manec_example)
```

---

rhat.bayesianecfit      *rhat.bayesianecfit*

---

**Description**

rhat.bayesianecfit

**Usage**

```
## S3 method for class 'bayesianecfit'
rhat(object, rhat_cutoff = 1.05, ...)
```

**Arguments**

object	An object of class <a href="#">bayesianecfit</a> as returned by <a href="#">bnec</a> .
rhat_cutoff	A <a href="#">numeric</a> vector indicating the Rhat cut-off used to test for model convergence.
...	Unused.

**Value**

A [list](#) containing a vector or Rhat values returned for each parameter for a [brmsfit](#) object, for each of the fitted models.

---

rhat.bayesianecfit      *rhat.bayesianecfit*

---

**Description**

rhat.bayesianecfit

**Usage**

```
## S3 method for class 'bayesianecfit'
rhat(object, ...)
```

**Arguments**

object	An object of class <a href="#">bayesianecfit</a> as returned by <a href="#">bnec</a> .
...	Unused.

**Value**

A named [numeric](#) vector containing Rhat values as returned for a [brmsfit](#) object for each of the estimated parameters.

---

sample_priors	<i>sample_priors</i>
---------------	----------------------

---

**Description**

Creates list or generates a plot of prior samples

**Usage**

```
sample_priors(priors, n_samples = 10000, plot = "ggplot")
```

**Arguments**

priors	An object of class <a href="#">brmsprior</a> from package <b>brms</b> .
n_samples	The number of prior samples to return.
plot	NA returns a <a href="#">list</a> of numeric vectors of sampled priors, "ggplot" (default) returns a <a href="#">ggplot</a> and "base" returns a histogram in base R.

**Value**

A [list](#) containing the initialisation values.

**See Also**

[bnec](#)

---

show_params	<i>show_params</i>
-------------	--------------------

---

**Description**

Displays non-linear equation and parameter names

**Usage**

```
show_params(model = "all")
```

**Arguments**

model	Removed in version 2.0. Use formula instead. Used to be a <a href="#">character</a> vector indicating the model(s) to fit. See Details for more information.
-------	--

**Value**

An [list](#) of [brmsformula](#).



**Examples**

```
library(bayesnec)
# default to all models (i.e. model = "all")
show_params()
# single model
show_params(model = "nec3param")
# group of models
show_params(model = c("nec3param", "ecx"))
```

---

summary.bayesmanecfit *summary.bayesmanecfit*

---

**Description**

summary.bayesmanecfit

**Usage**

```
## S3 method for class 'bayesmanecfit'
summary(object, ..., ecx = FALSE, ecx_vals = c(10, 50, 90))
```

**Arguments**

object	An object of class <a href="#">bayesmanecfit</a> as returned by <a href="#">bnec</a> .
...	Unused.
ecx	Should summary ECx values be calculated? Defaults to FALSE.
ecx_vals	ECx targets (between 1 and 99). Only relevant if ecx = TRUE. If no value is specified by the user, returns calculations for EC10, EC50, and EC90.

**Value**

A [list](#) containing a summary of the model fit as returned by a [brmsfit](#) object for each model.

---

summary.bayesnecfit *summary.bayesnecfit*

---

**Description**

summary.bayesnecfit

**Usage**

```
## S3 method for class 'bayesnecfit'
summary(object, ..., ecx = FALSE, ecx_vals = c(10, 50, 90))
```

**Arguments**

object	An object of class <code>bayesnecfit</code> as returned by <code>bneec</code> .
...	Unused.
ecx	Should summary ECx values be calculated? Defaults to FALSE.
ecx_vals	ECx targets (between 1 and 99). Only relevant if <code>ecx = TRUE</code> . If no value is specified by the user, returns calculations for EC10, EC50, and EC90.

**Value**

A summary of the fitted model as returned for a `brmsfit` object.

---

update.bnecfif	<i>Update an object of class <code>bnecfif</code> as fitted by function <code>bneec</code>.</i>
----------------	---

---

**Description**

Update an object of class `bnecfif` as fitted by function `bneec`.

**Usage**

```
## S3 method for class 'bnecfif'
update(
  object,
  newdata = NULL,
  recompile = NULL,
  x_range = NA,
  precision = 1000,
  sig_val = 0.01,
  loo_controls,
  force_fit = FALSE,
  ...
)
```

**Arguments**

object	An object of class <code>bnecfif</code> as fitted by function <code>bneec</code> .
newdata	Optional <code>data.frame</code> to update the model with new data. Data-dependent default priors will not be updated automatically.
recompile	A <code>logical</code> , indicating whether the Stan model should be recompiled. If NULL (the default), update tries to figure out internally, if recompilation is necessary. Setting it to FALSE will cause all Stan code changing arguments to be ignored.
x_range	A range of predictor values over which to consider extracting ECx.
precision	The length of the predictor vector used for posterior predictions, and over which to extract ECx values. Large values will be slower but more precise.

sig_val	Probability value to use as the lower quantile to test significance of the predicted posterior values against the lowest observed concentration (assumed to be the control), to estimate NEC as an interpolated NOEC value from smooth ECx curves.
loo_controls	A named <code>list</code> of two elements ("fitting" and/or "weights"), each being a named <code>list</code> containing the desired arguments to be passed on to <code>loo</code> (via "fitting") or to <code>loo_model_weights</code> (via "weights"). If "fitting" is provided with argument <code>pointwise = TRUE</code> (due to memory issues) and <code>family = "beta_binomial2"</code> , the <code>bnecfi</code> will fail because that is a custom family. If "weights" is not provided by the user, <code>bnecfi</code> will set the default method argument in <code>loo_model_weights</code> to "pseudobma". See <code>?loo_model_weights</code> for further info.
force_fit	Should model truly be updated in case either newdata of a new family is provided?
...	Further arguments to <code>brm</code> .

### Value

An object of class `bnecfi`. If one single model is returned, then also an object of class `bayesnecfi`; otherwise, if multiple models are returned, also an object of class `bayesmanecfi`.

### Examples

```
library(bayesnecfi)
data(manec_example)
# due to package size issues, `manec_example` does not contain original
# stanfit DSO, so need to recompile here
smaller_manec <- update(manec_example, chains = 1, iter = 50,
                       recompile = TRUE)
# original `manec_example` is fit with a Gaussian
# change to Beta distribution by adding newdata with original `nec_data$y`
# function will throw informative message.
beta_manec <- update(manec_example, newdata = nec_data, recompile = TRUE,
                    chains = 1, iter = 50, family = Beta(link = "identity"),
                    force_fit = TRUE)
```

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