

# Package ‘tmbstan’

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**Type** Package

**Title** MCMC Sampling from 'TMB' Model Object using 'Stan'

**Version** 1.0.0

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**Description** Enables all 'rstan' functionality for a 'TMB' model object, in particular MCMC sampling and chain visualization. Sampling can be performed with or without Laplace approximation for the random effects.

**Imports** methods, Rcpp

**Depends** rstan, TMB (>= 1.7.12)

**LinkingTo** StanHeaders, rstan, BH, Rcpp, RcppEigen, TMB

**License** GPL (>= 3)

**BugReports** <https://github.com/kaskr/tmbstan/issues>

**NeedsCompilation** yes

**RoxygenNote** 5.0.1

**Repository** CRAN

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tmbstan

*Draw MCMC samples from a TMB model object using Stan*


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## Description

Draw MCMC samples from a TMB model object using Stan

## Usage

```
tmbstan(obj, ..., lower = numeric(0), upper = numeric(0), laplace = FALSE,
        silent = TRUE, debug = FALSE)
```

## Arguments

obj	TMB model object.
...	Passed to <code>rstan::sampling</code> with some modifications - see details.
lower	Vector of lower parameter bounds.
upper	Vector of upper parameter bounds.
laplace	Apply the Laplace approximation to random subset of parameters ? The default disables the Laplace approximation.
silent	Be silent during sampling ?
debug	Should not be used.

## Details

tmbstan works for models with or without random effects.

By default a full Bayesian analysis is carried out, i.e. both parameters and random effects are sampled using MCMC. Models with random effects will thus have the Laplace approximation disabled. It is possible to mix the Laplace approximation with MCMC by setting `laplace=TRUE`. All methods provided by the `rstan` package can be applied to a fitted object. Get a complete list using `methods(class="stanfit")`.

Lower and upper bounds can be set using `lower` and `upper`. The bounds can be specified in one of two ways. Either in short format, i.e. have the same length as `obj$par`. Remaining parameters (the random effects) are set as unbounded in this case. Otherwise the bounds must be in long format, i.e. have the same length as the full parameter vector `obj$env$par` including the random effects. In both cases `-Inf` and `Inf` are valid components of `lower` and `upper` respectively. Note that initial values must be within the specified bounds.

The function arguments `...` are passed to `rstan`'s fitting function, see `?rstan::sampling`. A few notable arguments are:

- `chains` The number of chains.
- `iter` The number of iterations.
- `init` Initial values for the sampler. Behaves like `rstan` with some additions:
  - Default is "random" - see `?stan`.

- Special values  $\emptyset$  and "0" are allowed - see ?stan.
  - Additional special characters "par" and "last.par.best" are allowed and will be looked up in the TMB model object. The value "par" signifies to start from the defaults of the model object. If an optimization has been carried out, the initial value "last.par.best" will start from the MLE.
  - We also allow to pass a single numeric vector, or a list of numeric vectors. List length must match the number of chains. Vector lengths must match the number of sampled parameters. Names are currently ignored.
  - Parameters that do not follow the previous scheme (e.g. characters) are passed on to rstan unchanged. If in doubt, use `rstan::get_inits` to inspect the applied initial values.
- seed Random seed.

## Value

Object of class stanfit

## Examples

```
runExample("simple")
fit <- tmbstan(obj, chains=1)
class(fit) ## "stanfit"

## The available methods are
methods(class="stanfit")

## Not run:
## Pairs plot
pairs(fit, pars=names(obj$par))

## End(Not run)

## Trace plot
traceplot(fit, pars=names(obj$par), inc_warmup=TRUE)
```

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