# Package 'nof1gen'

November 18, 2019

Type Package
Title N-of-1 study general analysis tool
Version 0.5.0
<b>Depends</b> R (>= 2.10)
Imports rjags (>= 4-6), splines, combinat, MASS, jsonlite, ggplot2, scales, coda (>= 0.13)
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<b>Description</b> A package for running analysis for N of 1 studies. Runs Bayesian linear regression, ordinal/logistic regression, and poisson regression. Includes different plots to visualize the results.
License GPL-3
Encoding UTF-8
LazyData true
RoxygenNote 6.1.1
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nof1-package

mcnet: A package for N of 1 study analysis using Bayesian methods

## **Description**

A package for running N of 1 study trials

#### **Details**

An N of 1 trial is a clinical trial in which a single patient is the entire trial, a single case study. The main purpose of this package was to serve as an analysis tool for one of the PCORI grants we were working with. It is designed for N of 1 trials and can fit bayesian versions of linear regression, logistic/ordinal regression, and poisson regression. Package includes number of different plotting tools for visualization.

frequency\_plot

Frequency plot for raw data

# Description

Frequency plot for raw data

#### Usage

```
frequency_plot(nof1, ...)
```

#### **Arguments**

```
nof1 nof1 object created using nof1.data
```

... parameters to pass to geom\_bar or geom\_histogram.

# Examples

```
Y <- laughter$Y
Treat <- laughter$Treat
nof1 <- nof1.data(Y, Treat, ncat = 11, baseline = "Usual Routine", response = "ordinal")
frequency_plot(nof1)</pre>
```

kernel\_plot 3

#### **Description**

Creates a kernel density estimation plot for a specific outcome

#### Usage

```
kernel_plot(result, comp = T, ...)
```

#### **Arguments**

An object with information about the simulation. The object is derived from the output of nof1.run

... parameters to pass to geom\_histogram.

nof1.data Make an N of 1 object containing data, priors, and a jags model file

#### **Description**

Make an N of 1 object containing data, priors, and a jags model file

#### Usage

```
nof1.data(Y, Treat, response = NULL, ncat = NULL, bs.trend = F,
  y.time = NULL, knots.bt.block = NULL, block.no = NULL,
  corr.y = F, alpha.prior = NULL, beta.prior = NULL,
  eta.prior = NULL, dc.prior = NULL, c1.prior = NULL,
  rho.prior = NULL, hy.prior = NULL)
```

#### **Arguments**

Υ	Outcome of the study. This should be a vector with NA's included in time order.
Treat	Treatment indicator vector with same length as the outcome.
response	Type of outcome. Can be normal, binomial, poisson or ordinal.
ncat	Number of categories. Used in ordinal models.
bs.trend	Indicator for whether the model should adjust for trend using splines. The default is F.
y.time	parameter used for modeling splines. Time on the original scale with the same length as the outcome. Still under development.
knots.bt.block	parameter used for modeling splines. Currently, program only supports knots at the end of each block except for the last block if T.
block.no	parameter used for modeling splines. Block number with the same length as the outcome.
corr.y	Indicator for whether the model should model correlation. The default is F.

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alpha.prior Prior for the intercept of the model. beta.prior Prior for the treatment coefficient.

eta.prior Prior for modelling splines. Still under development.

dc.prior Prior for the length between cutpoints. Used only for ordinal logistic models.

c1.prior Prior for the first cutpoint. Used only for ordinal logistic models.

rho.prior Prior for the correlated error model. Still under development.

hy.prior Prior for the heterogeneity parameter. Supports uniform, gamma, and half nor-

mal for normal and binomial response and wishart for multinomial response. It should be a list of length 3, where first element should be the distribution (one of dunif, dgamma, dhnorm, dwish) and the next two are the parameters associated with the distribution. For example, list("dunif", 0, 5) give uniform prior with lower bound 0 and upper bound 5 for the heterogeneity parameter. For wishart distribution, the last two parameter would be the scale matrix and the degrees of

freedom.

#### Value

Creates list of variables that are used to run the model using nof1.run

Y Outcome Treat Treatment

ncat Number of categories for ordinal response
nobs Total number of observations in a study
Treat.name Treatment name besides baseline treatment

response The type of response variable

priors Priors that the code will be using. Default priors are used if prior was not speci-

fied

code Rjags model file code that is generated using information provided by the user.

To view model file inside R, use cat(nof1\$code).

#### **Examples**

```
###Blocker data example
laughter
Y <- laughter$Y
Treat <- laughter$Treat
nof1 <- nof1.data(Y, Treat, ncat = 11, response = "ordinal")
str(nof1)
cat(nof1$code)</pre>
```

nof1.normal.simulation

Normal data simulation

#### **Description**

Simulating sample normal data

nof1.run 5

#### Usage

```
nof1.normal.simulation(Base.size = 2, Treat.size = 8, prec = 0.5,
  alpha = 50, beta_A = -3, beta_B = -1)
```

nof1.run

Run the model using the nof1 object

## Description

This is the core function that runs the model in our program. Before running this function, we need to specify data, prior, JAGS code, etc. using nof1.data.

## Usage

```
nof1.run(nof1, inits = NULL, n.chains = 3, max.run = 1e+05,
  setsize = 10000, n.run = 50000, conv.limit = 1.05,
  extra.pars.save = NULL)
```

## **Arguments**

nof1	nof1 object created from nof1.data function	
inits	Initial values for the parameters being sampled. If left unspecified, program will generate reasonable initial values.	
n.chains	Number of chains to run	
max.run	Maximum number of iterations that user is willing to run. If the algorithm is not converging, it will run up to max. run iterations before printing a message that it did not converge	
setsize	Number of iterations that are run between convergence checks. If the algorithm converges fast, user wouldn't need a big setsize. The number that is printed between each convergence checks is the gelman-rubin diagnostics and we would want that to be below the conv.limit the user specifies.	
n.run	Final number of iterations that the user wants to store. If after the algorithm converges, user wants less number of iterations, we thin the sequence. If the user wants more iterations, we run extra iterations to reach the specified number of runs	
conv.limit	Convergence limit for Gelman and Rubin's convergence diagnostic.	
extra.pars.save		
	Parameters that user wants to save besides the default parameters saved. See	

# Value

nof1	nof1 object
inits	Initial values that are either specified by the user or generated as a default
pars.save	Parameters that are saved. Add more parameters in extra.pars.save if other variables are desired
data_rjags	Data that is put into rjags function jags.model
burnin	Half of the converged sequence is thrown out as a burnin

code using cat(nof1\$code) to see which parameters can be saved.

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n.thin If the number of iterations user wants (n.run) is less than the number of converged sequence after burnin, we thin the sequence and store the thinning inter-

val

samples MCMC samples stored using jags. The returned samples have the form of

mcmc.list and can be directly applied to coda functions

max.gelman Maximum Gelman and Rubin's convergence diagnostic calculated for the final

sample

# **Examples**

```
laughter
Y <- laughter$Y
Treat <- laughter$Treat
nof1 <- nof1.data(Y, Treat, ncat = 11, baseline = "Usual Routine", response = "ordinal")
result <- nof1.run(nof1)
summary(result$samples)</pre>
```

raw\_table

Summary data table for nof1

#### **Description**

Provides a summary data table for the particular outcome in a particular dataset.

#### Usage

```
raw_table(nof1)
```

#### **Arguments**

nof1 nof1 object created using nof1.data

#### Value

Gives a comprhensive table with several statistical values. Each column indicates the value given. For a normal or poisson response type the following are given:

Treat	The treatment recieved
mean	The average value of the outcome
sd	The standard deviation for the outcome
2.5%	2.5% of the data are equal to or less than this value
50%	50% of the data are equal to or less than this value
97.5%	97.5% of the data are equal to or less than this value

For a binomial or ordinal response type, returns a table where first row is each treatment and the following rows are the the number of data points taken at each possible value.

#### **Examples**

```
Y <- laughter$Y
Treat <- laughter$Treat
nof1 <- nof1.data(Y, Treat, ncat = 11, baseline = "Usual Routine", response = "ordinal")
raw_table(nof1)</pre>
```

```
stacked_percent_barplot
```

Stacked\_percent\_barplot for raw data (for ordinal or binomial data)

#### **Description**

Stacked\_percent\_barplot for raw data (for ordinal or binomial data)

## Usage

```
stacked_percent_barplot(nof1)
```

#### **Arguments**

nof1

nof1 object created using nof1.data

#### **Examples**

```
Y <- laughter$Y
Treat <- laughter$Treat
nof1 <- nof1.data(Y, Treat, ncat = 11, baseline = "Usual Routine", response = "ordinal")
stacked_percent_barplot(nof1)</pre>
```

summarize\_nof1

Function to present a summary of our results

#### **Description**

A neat function to summarize the results.

#### Usage

```
summarize_nof1(result, alpha = 0.05)
```

#### **Arguments**

result A list which contains data file created by nof1.data and the result file created

by nof1.run

alpha The alpha value for the confidence interval. If no value is entered will give the

95% confidence interval.

#### Value

The function computes and returns a list of summary statistics of the raw data and the fitted model.

raw.y.mean The raw mean of the outcome for each treatment raw.y.median The raw median of the outcome for each treatment post.coef.mean The posterior mean of the coefficient for each treatment post.coef.median

The posterior median of the coefficient for each treatment

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```
post.y.mean
                  The posterior mean of the outcome for each treatment
post.y.median
                  The posterior median of the outcome for each treatment
                  The credible interval of the coefficient for each treatment
post.coef.ci
                  The credible interval of the outcome for each treatment
post.y.ci
comp.treat.post.coef
                  The posterior quantiles of one coefficient minus the other when comparing two
                  treatments
comp.treat.post.y
                  The posterior quantiles of outcome minus the other when comparing two treat-
                  ments
p.comp.coef.greater.0
```

The posterior probability of one coefficient minus the other greater than 0

time\_series\_plot

time series plot across different interventions

#### **Description**

time series plot across different interventions

#### Usage

```
time_series_plot(result, overlay.with.model = F, plot.by.treat = T,
  trial.start = NULL, trial.end = NULL, timestamp.format = NULL)
```

#### **Arguments**

trial.start Start time of the trial specified with timestamp.format. End time of the trial specified with timestamp.format. trial.end timestamp.format Format of the trial.start and trial.end. nof1 object created using nof1.data nof1

## **Examples**

```
Y <- laughter$Y
Treat <- laughter$Treat</pre>
nof1 <- nof1.data(Y, Treat, ncat = 11, baseline = "Usual Routine", response = "ordinal")</pre>
timestamp <- seq(as.Date('2015-01-01'),as.Date('2016-01-31'), length.out = length(Y))
time_series_plot(nof1, timestamp = timestamp, timestamp.format = "%m-%d-%Y", Outcome.name = "Stress")
```

wrap

For PCORI purposes

#### **Description**

For PCORI purposes

# Usage

```
wrap(data, metadata)
```

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wrap2

For PCORI purposes

# Description

For PCORI purposes

# Usage

wrap2(data, metadata)

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