# Package 'GLMMcosinor'

November 1, 2024

Type Package

```
Title Fit a Cosinor Model Using a Generalized Mixed Modeling Framework
Version 0.2.1
Description Allows users to fit a cosinor model using the 'glmmTMB' framework.
      This extends on existing cosinor modeling packages, including 'cosinor'
      and 'circacompare', by including a wide range of available link functions
      and the capability to fit mixed models. The cosinor model is described by
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2 amp\_acro

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# **Contents**

	amp_acro	2
	autoplot.cglmm	
	cglmm	6
	cosinor_mixed	7
	fit_model_and_process	8
	polar_plot	9
	polar_plot.cglmm	11
	predict.cglmm	14
	print.cglmm	14
	print.cglmmSubTest	15
	print.cglmmSummary	16
	print.cglmmTest	17
	sigma.cglmm	18
	simulate_cosinor	19
	summary.cglmm	20
	test_cosinor_components	21
	test_cosinor_levels	23
	update_formula_and_data	24
	vitamind	26
Index		27

amp\_acro

Used to specify a cosinor component in the model formula.

# Description

Checks the validity of user inputs before creating an updated formula and associated modifications to the data.frame.

#### Usage

```
amp_acro(time_col, n_components = 1, group, period, ...)
```

amp\_acro 3

#### **Arguments**

time\_col A numeric column within the data.frame() passed by via the data arg containing the time values.

n\_components The Number of cosinor components in the model.

group A vector of the names for the group factors (column names within the data.frame() passed by via the data arg).

period A numeric value or vector containing the period. The number of values should be equal to n\_components.

... Extra arguments for use within GLMMcosinor.

#### Value

A data.frame and formula appropriate for use by data\_processor().

```
# Single component cosinor model
cglmm(
 vit_d ~ amp_acro(time_col = time, group = "X", period = 12),
 data = vitamind
)
# 2-component cosinor model with simulated data
sim_data <- simulate_cosinor(</pre>
 n = 500,
 mesor = 5,
 amp = c(2, 1),
 acro = c(1, 1.5),
 beta.mesor = 2,
 beta.amp = c(2, 1),
 beta.acro = c(1, 1.5),
 family = "gaussian",
 period = c(12, 6),
 n_{components} = 2,
 beta.group = TRUE,
)
cglmm(
 Y ~ group + amp_acro(times,
   n_{components} = 2,
   group = "group",
   period = c(12, 6)
 ),
 data = sim_data,
 family = gaussian
)
```

4 autoplot.cglmm

autoplot.cglmm

Plot a cosinor model

#### **Description**

Given a cglmm model fit, generate a plot of the data with the fitted values. Optionally allows for plotting by covariates

#### Usage

```
## S3 method for class 'cglmm'
autoplot(
  object,
 ci_level = 0.95,
  x_str,
  type = "response",
  xlims,
  pred.length.out,
  points_per_min_cycle_length = 20,
  superimpose.data = FALSE,
  data_opacity = 0.3,
  predict.ribbon = TRUE,
  ranef_plot = NULL,
  cov_list = NULL,
  quietly = TRUE,
)
```

#### **Arguments**

object An cglmm object.

ci\_level The level for calculated confidence intervals. Defaults to 0.95.

x\_str A character vector naming variable(s) to be plotted. Default has no value and

plots all groups.

type A character that will be passed as an argument to predict.cglmm(), specify-

ing the type of prediction (e.g, "response", or "link"). See ?glmmTMB::predict.glmmTMB

for full list of possible inputs.

xlims A vector of length two containing the limits for the x-axis.

pred.length.out

An integer value that specifies the number of predicted data points. The larger

the value, the more smooth the fitted line will appear. If missing, uses  $points\_per\_min\_cycle\_length$ 

to generate a sensible default value.

points\_per\_min\_cycle\_length

Used to determine the number of samples to create plot if pred.length.out is

missing.

autoplot.cglmm 5

superimpose.data

A logical. If TRUE, data from the original data used to fit the model (object)

will be superimposed over the predicted fit.

data\_opacity A number between 0 and 1 inclusive that controls the opacity of the superim-

posed data. (Used as the alpha when calling ggplot2::geom\_point()).

predict.ribbon A logical. If TRUE, a prediction interval is plotted.

ranef\_plot Specify the random effects variables that you wish to plot. If not specified, only

the fixed effects will be visualized.

cov\_list Specify the levels of the covariates that you wish to plot as a list. For ex-

ample, if you have two covariates: var1, and var 2, you could fix the level to be plotted as such cov\_list = list(var1 = 'a', var2 = 1), where 'a' is a level in 'var1', and 1 is a level in 'var2'. See the examples for a demonstration. If not specified, the reference level of the covariate(s) will be used. points\_per\_min\_cycle\_length is the number of points plotted per the mini-

mum cycle length (period) of all cosinor components in the model.

quietly A logical. If TRUE, shows warning messages when wrangling data and fitting

model. Defaults to TRUE.

... Additional, ignored arguments.

#### Value

Returns a ggplot object.

```
# A simple model
model <- cglmm(</pre>
 vit_d ~ X + amp_acro(time, group = "X", period = 12),
 data = vitamind
autoplot(model, x_str = "X")
# Plotting a model with various covariates
test_data <- vitamind[vitamind$X == 1, ]</pre>
test_data$var1 <- sample(c("a", "b", "c"), size = nrow(test_data), replace = TRUE)
test_data$var2 <- rnorm(n = nrow(test_data))</pre>
object <- cglmm(</pre>
 vit_d ~ amp_acro(time, period = 12) + var1 + var2,
 data = test_data
autoplot(object,
 cov_list = list(
   var1 = "a",
   var2 = 1
 ),
 superimpose.data = TRUE
)
```

6 cglmm

cglmm	Fit cosinor model with {glmmTMB}	
-------	----------------------------------	--

# Description

Given an outcome and time variable, fit the cosinor model with optional covariate effects.

## Usage

```
cglmm(
  formula,
  data,
  family = stats::gaussian(),
  quietly = TRUE,
  dispformula = ~1,
  ziformula = ~0,
  ...
)
```

## Arguments

formula	A formula specifying the cosinor model to be fit. The cosinor portion of the formula is controlled by including amp_acro() on the right hand side of the formula. See amp_acro for more details.
data	A data.frame containing the variables used in the model.
family	A family function or a character string naming a family function. See ?family and ?glmmTMB::family_glmmTMB for options.
quietly	A logical. If TRUE, shows warning messages when wrangling data and fitting model. Defaults to TRUE.
dispformula	A one-sided (i.e., no response variable) formula for dispersion combining fixed and random effects, including cosinor components using $amp\_acro()$ . Defaults to $\sim$ 1.
ziformula	A one-sided (i.e., no response variable) formula for zero-inflation combining fixed and random effects, including cosinor components using $amp\_acro()$ . Defaults to $\sim 0$ .
	Optional additional arguments passed to glmmTMB::glmmTMB().

#### Value

Returns a fitted cosinor model as a cglmm object.

#### References

Tong, YL. Parameter Estimation in Studying Circadian Rhythms, Biometrics (1976). 32(1):85–94.

cosinor\_mixed 7

#### **Examples**

```
# Single component cosinor model
 vit_d ~ amp_acro(time_col = time, group = "X", period = 12),
 data = vitamind
)
# 2-component cosinor model with simulated data
sim_data <- simulate_cosinor(</pre>
 n = 500,
 mesor = 5,
 amp = c(2, 1),
 acro = c(1, 1.5),
 beta.mesor = 2,
 beta.amp = c(2, 1),
 beta.acro = c(1, 1.5),
 family = "gaussian",
 period = c(12, 6),
 n_{components} = 2,
 beta.group = TRUE,
)
cglmm(
 Y ~ group + amp_acro(times,
   n_{components} = 2,
   group = "group",
   period = c(12, 6)
 ),
 data = sim_data,
 family = gaussian
)
```

cosinor\_mixed

cosinor\_mixed dataset for cosinor modeling examples.

#### **Description**

Simulated data set to illustrate a mixed cosinor model. The Y column contains a simulated outcome variable that varies over the time variable (times). The subject column is a grouping variable that can be used as a random effect. The rhythm has a period of 24 hours. Data was simulated using simulate\_cosinor.

#### Usage

cosinor\_mixed

## **Format**

A data.frame with 3 variables: Y, times, and subject.

#### **Description**

Fit the cosinor GLMM model using the output from update\_formula\_and\_data() and a new formula

#### **Usage**

```
fit_model_and_process(obj, formula, ...)
```

#### **Arguments**

obj Output from update\_formula\_and\_data().

formula A (optionally) new formula to use when fitting the cosinor model (maybe with random effects) or other covariates found in the data.

Optional additional arguments passed to glmmTMB::glmmTMB().

#### Value

Returns a fitted cosinor model as a cglmm object.

```
# Use vitamind data but add a "patient" identifier used as a random effect
vitamind2 <- vitamind</pre>
vitamind2$patient <- sample(</pre>
 LETTERS[1:5],
 size = nrow(vitamind2), replace = TRUE
# Use update_formula_and_data() to perform wrangling steps of cglmm()
# without yet fitting the model
data_and_formula <- update_formula_and_data(</pre>
 data = vitamind2,
 formula = vit_d ~ X + amp_acro(time,
    group = "X",
   period = 12
 )
)
# print formula from above
data_and_formula$newformula
# fit model while adding random effect to cosinor model formula.
mod <- fit_model_and_process(</pre>
 obj = data_and_formula,
```

polar\_plot 9

```
formula = update.formula(
    data_and_formula$newformula, . ~ . + (1 | patient)
)

mod
mod$fit # printing the `glmmTMB` model within shows Std.Dev. of random effect
```

polar\_plot

Generates a polar plot with elliptical confidence intervals

#### **Description**

Generates a polar plot with elliptical confidence intervals

#### Usage

```
polar_plot(
  ci_level = 0.95,
  n_breaks = 5,
  component_index = NULL,
  grid_angle_segments = 8,
  radial_units = c("radians", "degrees", "period"),
  clockwise = FALSE,
  text\_size = 3,
  text_opacity = 0.5,
  fill_colors,
  ellipse_opacity = 0.3,
  circle_linetype = "dotted",
  start = c("right", "left", "top", "bottom"),
  view = c("full", "zoom", "zoom_origin"),
  overlay_parameter_info = FALSE,
  quietly = TRUE,
  show_component_labels = TRUE,
  xlims,
  ylims,
)
```

#### Arguments

x An object of class cglmm

ci\_level The level for calculated confidence ellipses. Defaults to 0.95.

n\_breaks The number of concentric circles that will be plotted using the scales::breaks\_pretty()

function. By default, 5 breaks will be used. The number of breaks may be adjusted to result in an even interval. For example, if n\_breaks is 3, but the

10 polar\_plot

maximum plot radius is 8, instead of plotting circles in intervals in 1.6, this interval will be rounded to 2 to result in the sequence: 0, 2, 4, 6, 8. See ?scales::breaks\_pretty for more details.

#### component\_index

A number that corresponds to a particular component from the cglmm() object that will be used to create polar plot. If missing (default), then plots for all components will be arranged in the returned plot. If a single or multiple values are provided, then these components will be returned. (for example component\_index = 1, component\_index = c(1, 3)).

#### grid\_angle\_segments

An integer. Determines the total number of segments in the background of the polar plot. For example, a value of 4 will create quadrants around the origin. Defaults to 8.

radial\_units

A character specifying the angular units of the plot. Possible values are one of c('radians', 'degrees', 'period'). These units relate to the period of the component being visualized.

'radians':  $[0, 2\pi]$ 'degrees': [0, 360]'period': [0, period]

clockwise

A logical. If TRUE, the angles increase in a clockwise fashion. If FALSE, anti-clockwise. Defaults to FALSE.

text\_size

A number controlling the font size of the text labels. Defaults to 3.

text\_opacity
fill\_colors

A character vector containing colors that will be mapped to levels within a group. If the model has components with different number of levels per factor, the length of this input should match the greatest number of levels. If not, or if the number of levels exceeds the length of the default argument (8), colors are

A numeric between 0 and 1 inclusive that controls the opacity of the text labels.

generated using rainbow().

ellipse\_opacity

A numeric between 0 and 1 inclusive that controls the opacity of the confidence ellipses. Defaults to 0.3.

circle\_linetype

A character or numeric that determines the linetype of the radial circles in background of the polar plot. See ?linetype for more details.

start

A character, within c("right", "left", "top", "bottom") that determines where angle 0 is located. If start = "top", and clockwise = TRUE, the angle will rotate clockwise, starting at the '12 o-clock' position on a clock.

view

A character, within c("full", "zoom", "zoom\_origin") that controls the view of the plots.

'full': maintains a full view of the polar plot, including the background radial circles.

'zoom': finds the minimum view window which contains all confidence ellipses.

<sup>&#</sup>x27;zoom\_origin': zooms into the confidence ellipses (like "zoom"), but also keeps the origin within f

11 polar\_plot.cglmm

```
overlay_parameter_info
                  A logical argument. If TRUE, more information about the acrophase and am-
                  plitude are displayed on the polar plots.
quietly
                  Analogous to verbose, this logical argument controls whether messages are
                  displayed in the console.
show_component_labels
                  Logical argument, TRUE by default. When TRUE, the polar plots have labels
                  corresponding to their components.
                  A vector of length two containing the limits for the x-axis.
xlims
ylims
                  A vector of length two containing the limits for the y-axis.
                  Additional, ignored arguments.
```

#### Value

. . .

Returns a ggplot object.

#### **Examples**

```
data(vitamind)
model <- cglmm(</pre>
  vit_d ~ X + amp_acro(time, group = "X", period = 12),
  data = vitamind
polar_plot(model, radial_units = "period")
```

polar\_plot.cglmm

Generates a polar plot with elliptical confidence intervals

#### **Description**

Generates a polar plot with elliptical confidence intervals

## Usage

```
## S3 method for class 'cglmm'
polar_plot(
  Х,
  ci_level = 0.95,
  n_breaks = 5,
  component_index = NULL,
  grid_angle_segments = 8,
  radial_units = c("radians", "degrees", "period"),
  clockwise = FALSE,
  text\_size = 3.5,
  text_opacity = 1,
  fill_colors,
  ellipse_opacity = 0.3,
```

12 polar\_plot.cglmm

```
circle_linetype = "dotted",
start = c("right", "left", "top", "bottom"),
view = c("full", "zoom", "zoom_origin"),
overlay_parameter_info = FALSE,
quietly = TRUE,
show_component_labels = TRUE,
xlims,
ylims,
...
```

#### Arguments

x An object of class cglmm

ci\_level The level for calculated confidence ellipses. Defaults to 0.95.

n\_breaks The number of concentric circles that will be plotted using the scales::breaks\_pretty()

function. By default, 5 breaks will be used. The number of breaks may be adjusted to result in an even interval. For example, if n\_breaks is 3, but the maximum plot radius is 8, instead of plotting circles in intervals in 1.6, this interval will be rounded to 2 to result in the sequence: 0, 2, 4, 6, 8. See ?scales::breaks\_pretty for more details.

component\_index

A number that corresponds to a particular component from the cglmm() object that will be used to create polar plot. If missing (default), then plots for all components will be arranged in the returned plot. If a single or multiple values are provided, then these components will be returned. (for example component\_index = 1, component\_index = c(1, 3)).

grid\_angle\_segments

An integer. Determines the total number of segments in the background of the polar plot. For example, a value of 4 will create quadrants around the origin. Defaults to 8.

radial\_units

A character specifying the angular units of the plot. Possible values are one of c('radians', 'degrees', 'period'). These units relate to the period of the component being visualized.

'radians':  $[0, 2\pi]$ 'degrees': [0, 360]'period': [0, period]

clockwise

A logical. If TRUE, the angles increase in a clockwise fashion. If FALSE, anti-clockwise. Defaults to FALSE.

text\_size

A number controlling the font size of the text labels. Defaults to 3.

text\_opacity

A numeric between 0 and 1 inclusive that controls the opacity of the text labels.

fill\_colors

A character vector containing colors that will be mapped to levels within a group. If the model has components with different number of levels per factor, the length of this input should match the greatest number of levels. If not, or if the number of levels exceeds the length of the default argument (8), colors are

generated using rainbow().

13 polar\_plot.cglmm

ellipse\_opacity

A numeric between 0 and 1 inclusive that controls the opacity of the confidence ellipses. Defaults to 0.3.

circle\_linetype

A character or numeric that determines the linetype of the radial circles in background of the polar plot. See ?linetype for more details.

start

A character, within c("right", "left", "top", "bottom") that determines where angle 0 is located. If start = "top", and clockwise = TRUE, the angle will rotate clockwise, starting at the '12 o-clock' position on a clock.

view

A character, within c("full", "zoom", "zoom\_origin") that controls the view of the plots.

'full': maintains a full view of the polar plot, including the background radial circles.

'zoom': finds the minimum view window which contains all confidence ellipses.

'zoom\_origin': zooms into the confidence ellipses (like "zoom"), but also keeps the origin within f

overlay\_parameter\_info

A logical argument. If TRUE, more information about the acrophase and amplitude are displayed on the polar plots.

quietly

Analogous to verbose, this logical argument controls whether messages are displayed in the console.

show\_component\_labels

Logical argument, TRUE by default. When TRUE, the polar plots have labels corresponding to their components.

xlims A vector of length two containing the limits for the x-axis.

A vector of length two containing the limits for the y-axis. ylims

Additional, ignored arguments.

#### Value

Returns a ggplot object.

```
model <- cglmm(</pre>
  vit_d ~ X + amp_acro(time, group = "X", period = 12),
  data = vitamind
polar_plot(model, radial_units = "period")
```

14 print.cglmm

predict.cglmm

Predict from a cosinor model

#### **Description**

Given a time variable and optional covariates, generate predicted values from a cosinor fit. Default prediction is the mean value, optionally can predict at a given month

#### Usage

```
## S3 method for class 'cglmm'
predict(object, newdata, ...)
```

# Arguments

```
object An object of class cglmm.

newdata Optional new data.

other arguments passed to glmmTMB:::predict.glmmTMB.
```

#### Value

Returns predicted values from the cosinor model.

## **Examples**

```
fit <- cglmm(vit_d ~ X + amp_acro(time,
   group = "X",
   n_components = 1,
   period = 12
), data = vitamind)
predict(fit)</pre>
```

print.cglmm

Print a brief summary of the cglmm model.

## **Description**

Print a brief summary of the cglmm model.

#### Usage

```
## S3 method for class 'cglmm'
print(x, digits = getOption("digits"), ...)
```

print.cglmmSubTest 15

## **Arguments**

```
    x A cglmm object.
    digits Controls the number of digits displayed in the summary output.
    ... Additional, ignored arguments.
```

#### Value

```
print(x) returns x invisibly.
```

### **Examples**

```
# Single component cosinor model
cglmm(
  vit_d ~ amp_acro(time_col = time, group = "X", period = 12),
  data = vitamind
)
```

 $\verb"print.cglmmSubTest"$ 

Print test of model

#### **Description**

Print test of model

#### Usage

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

#### **Arguments**

```
x A sub_test_cosinor object.... Additional, ignored arguments.
```

#### Value

```
print(x) returns x invisibly.
```

```
data_2_component <- simulate_cosinor(
  n = 10000,
  mesor = 5,
  amp = c(2, 5),
  acro = c(0, pi),
  beta.mesor = 4,
  beta.amp = c(3, 4),
  beta.acro = c(0, pi / 2),</pre>
```

```
family = "gaussian",
 n_{components} = 2,
 period = c(10, 12),
 beta.group = TRUE
)
mod_2_component <- cglmm(</pre>
 Y ~ group + amp_acro(times,
   n_components = 2, group = "group",
   period = c(10, 12)
 ),
 data = data_2_component
)
test_output <- test_cosinor_levels(</pre>
 mod_2_component,
 param = "amp",
 x_str = "group"
)
print(test_output$global.test)
```

print.cglmmSummary

Print the summary of a cosinor model

## **Description**

Print the summary of a cosinor model

#### Usage

```
## S3 method for class 'cglmmSummary'
print(x, digits = getOption("digits"), ...)
```

#### Arguments

```
x An object of class cglmmSummarydigits Controls the number of digits displayed in the summary output... Currently unused
```

#### Value

print returns x invisibly.

```
fit <- cglmm(vit_d ~ X + amp_acro(time,
  group = "X",
  n_components = 1,
  period = 12
), data = vitamind)
summary(fit)</pre>
```

print.cglmmTest 17

print.cglmmTest

Print results of test of cosinor model

# Description

Print results of test of cosinor model

#### Usage

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

# Arguments

```
x A test_cosinor object.
... Arguments passed to print
```

#### Value

```
print(x) returns x invisibly.
```

```
data_2_component <- simulate_cosinor(</pre>
  n = 10000,
  mesor = 5,
  amp = c(2, 5),
  acro = c(0, pi),
  beta.mesor = 4,
  beta.amp = c(3, 4),
  beta.acro = c(0, pi / 2),
  family = "gaussian",
  n_{components} = 2,
  period = c(10, 12),
  beta.group = TRUE
)
mod_2_component <- cglmm(</pre>
  Y ~ group + amp_acro(times,
    n_components = 2, group = "group",
    period = c(10, 12)
  ),
  data = data_2_component
test_cosinor_levels(
  mod_2_component,
  param = "amp",
  x_str = "group"
)
```

18 sigma.cglmm

sigma.cglmm

Extract residual standard deviation or dispersion parameter

## Description

```
see ?glmmTMB::sigma for more details.
```

#### Usage

```
## S3 method for class 'cglmm'
sigma(object, ...)
```

## Arguments

```
object An object of class cglmm.
... (ignored; for method compatibility)
```

#### Value

a numeric.

```
testdata_poisson <- simulate_cosinor(100,</pre>
 n_period = 2,
 mesor = 7,
 amp = c(0.1, 0.5),
 acro = c(1, 1),
 beta.mesor = 4.4,
 beta.amp = c(0.1, 0.46),
 beta.acro = c(0.5, -1.5),
 family = "poisson",
 period = c(12, 6),
 n_{components} = 2,
 beta.group = TRUE
)
mod <- cosinor_model <- cglmm(</pre>
 Y \sim group + amp_acro(times,
   period = c(12, 6),
   n_{components} = 2,
   group = "group"
 ),
 data = testdata_poisson,
 family = glmmTMB::nbinom1()
sigma(mod)
```

simulate\_cosinor 19

 $simulate\_cosinor$ 

Simulate data from a cosinor model

## Description

This function simulates data from a cosinor model with a single covariate, where the time scale is month, and optionally allows for single covariate effects on the mean, amplitude, and acrophase.

## Usage

```
simulate_cosinor(
    n,
    mesor,
    amp,
    acro,
    period = 24,
    n_components,
    beta.group = FALSE,
    beta.mesor,
    beta.amp,
    beta.acro,
    n_period = 1,
    family = c("gaussian", "poisson", "binomial", "gamma"),
    ...
)
```

#### **Arguments**

n	The sample size. An integer greater than 0.
mesor	A numeric. The MESOR (midline estimating statistic of rhythm) for group = 0. The MESOR is independent of the cosinor components, so only one value is allowed even if there are multiple components in the data being simulated.
атр	A numeric. The amplitude value (for group = 0 if grouped data are being simulated (beta.group = TRUE)). If simulating data with multiple components, specify a vector with values for each component. E.g. amp = $c(5, 10)$ .
acro	A numeric. The acrophase value in radians (for group = 0 if grouped data are being simulated (beta.group = TRUE)). If simulating data with multiple components, specify a vector with values for each component. E.g. acr = $c(0, pi)$ for two components.
period	The period of the rhythm data (for group = 0 if grouped data are being simulated (beta.group = TRUE)). If simulating data with multiple components, specify a vector with values for each component. E.g. period = $c(12, 6)$ for two components.
n_components	The number of components in the model. This must match the length of the inputs for amp and acro.

20 summary.cglmm

beta.group	A logical. If TRUE a second group of data will be simulated and included in the returned data set. If FALSE, beta.acro, beta.mesor, and beta.amp arguments will be ignored.
beta.mesor	A numeric. The MESOR value term for group = 1
beta.amp	A numeric. The amplitude value for group = 1. If simulating data with multiple components, specify a vector with values for each component. E.g. amp = $c(2, 8)$ .
beta.acro	A numeric. The acrophase value in radians (for group = 1. If simulating data with multiple components, specify a vector with values for each component. E.g. $acr = c(2, 5)$ for two components.
n_period	A numeric. The number of cycles of the rhythm to be simulated.
family	A character. The family (see ?family) of the simulated dataset. Can handle values in c("poisson", "binomial", "gamma", "gaussian").
	Extra arguments, including alpha that controls the shape argument when sampling from a gamma distribution (when family = "gamma"; default is 1), and sd (standard deviation) which is used when sampling from a normal distribution (when family = "gaussian"; default is 1). To specify these parameters for the beta (treatment) group, use beta.alpha and beta.sd

#### Value

Returns simulated data in a data. frame.

#### **Examples**

```
simulate_cosinor(
    n = 100,
    mesor = 1,
    amp = 1,
    acro = 1,
    period = 24,
    family = "gaussian"
)
```

summary.cglmm

Summarize a cosinor model Given a time variable and optional covariates, generate inference a cosinor fit. Gives estimates, confidence intervals, and tests for the raw parameters, and for the mean, amplitude, and acrophase parameters. If the model includes covariates, the function returns the estimates of the mean, amplitude, and acrophase for the group with covariates equal to 1 and equal to 0. This may not be the desired result for continuous covariates.

## **Description**

Summarize a cosinor model Given a time variable and optional covariates, generate inference a cosinor fit. Gives estimates, confidence intervals, and tests for the raw parameters, and for the mean, amplitude, and acrophase parameters. If the model includes covariates, the function returns the estimates of the mean, amplitude, and acrophase for the group with covariates equal to 1 and equal to 0. This may not be the desired result for continuous covariates.

#### Usage

```
## S3 method for class 'cglmm'
summary(object, ci_level = 0.95, ...)
```

#### **Arguments**

```
object An object of class cglmm

ci_level The level for calculated confidence intervals. Defaults to 0.95.

... Currently unused
```

#### Value

Returns a summary of the cglmm model as a cglmmSummary object.

#### **Examples**

```
fit <- cglmm(vit_d ~ X + amp_acro(time,
   group = "X",
   n_components = 1,
   period = 12
), data = vitamind)
summary(fit)</pre>
```

test\_cosinor\_components

Test for differences in a cosinor model between components.

## Description

Given a time variable and optional covariates, generate inference a cosinor fit. For the covariate named (or vector of covariates), this function performs a Wald test comparing the group with covariates equal to 1 to the group with covariates equal to 0. This may not be the desired result for continuous covariates.

## Usage

```
test_cosinor_components(
    x,
    x_str = NULL,
    param = "amp",
    comparison_A = 1,
    comparison_B = 2,
    level_index = 0,
    ci_level = 0.95
)
```

## Arguments

x	An cglmm object.
x_str	A character. The name of the grouping variable within which differences in the selected cosinor characteristic (amplitude or acrophase) will be tested. If there is no grouping variable in the model, then this can be left as NULL (default).
param	A character. Either "amp" or "acr" for testing differences in amplitude or acrophase, respectively.
comparison_A	An integer. Refers to the component number that is to act as the reference group. for the comparison.
comparison_B	An integer. Refers to the component number that is to act as the comparator group
level_index	An integer. If comparison_type = "components", level_index indicates which level of the grouping variable is being used for the comparison between components.
ci_level	The level for calculated confidence intervals. Defaults to 0.95.

## Value

Returns a test\_cosinor object.

```
data_2_component <- simulate_cosinor(
  n = 10000,
  mesor = 5,
  amp = c(2, 5),
  acro = c(0, pi),
  beta.mesor = 4,
  beta.amp = c(3, 4),
  beta.acro = c(0, pi / 2),
  family = "gaussian",
  n_components = 2,
  period = c(10, 12),
  beta.group = TRUE
)</pre>
```

test\_cosinor\_levels 23

test\_cosinor\_levels

Test for differences in a cosinor model between levels of the grouping variable.

## Description

Given a time variable and optional covariates, generate inference a cosinor fit. For the covariate named (or vector of covariates), this function performs a Wald test comparing the group with covariates equal to 1 to the group with covariates equal to 0. This may not be the desired result for continuous covariates.

#### Usage

```
test_cosinor_levels(
    x,
    x_str,
    param = "amp",
    comparison_A,
    comparison_B,
    component_index = 1,
    ci_level = 0.95
)
```

## Arguments

X	An cglmm object.
x_str	A character. The name of the grouping variable within which differences in the selected cosinor characteristic (amplitude or acrophase) will be tested.
param	A character. Either "amp" or "acr" for testing differences in amplitude or acrophase, respectively.
comparison_A	An integer, or string. Refers to the first level within the grouping variable x_str that is to act as the reference group in the comparison. Ensure that it corresponds to the name of the level in the original dataset.
comparison_B	An integer, or string. Refers to the second level within the grouping variable x_str that is to act as the comparator group in the comparison. Ensure that it corresponds to the name of the level in the original dataset.

```
component_index
```

An integer. If comparison\_type = "levels", component\_index indicates which component is being compared between the levels of the grouping variable.

ci\_level

The level for calculated confidence intervals. Defaults to 0.95.

#### Value

Returns a test\_cosinor object.

## **Examples**

```
data_2_component <- simulate_cosinor(</pre>
  n = 10000,
  mesor = 5,
  amp = c(2, 5),
  acro = c(0, pi),
  beta.mesor = 4,
  beta.amp = c(3, 4),
  beta.acro = c(0, pi / 2),
  family = "gaussian",
  n_{components} = 2,
  period = c(10, 12),
  beta.group = TRUE
mod_2_component <- cglmm(</pre>
  Y ~ group + amp_acro(times,
    n_components = 2, group = "group",
    period = c(10, 12)
  ),
  data = data_2_component
)
test_cosinor_levels(mod_2_component, param = "amp", x_str = "group")
```

update\_formula\_and\_data

Update data and formula for fitting cglmm model

#### **Description**

Update data and formula for fitting cglmm model

#### Usage

```
update_formula_and_data(
  data,
  formula,
  family = "gaussian",
  quietly = TRUE,
  dispformula = ~1,
```

```
ziformula = ~0
)
```

#### **Arguments**

input data for fitting cglmm model.

formula model formula, specified by user including amp\_acro().

family the model family.

quietly controls whether messages from amp\_acro are displayed. TRUE by default dispformula

The formula specifying the dispersion model

ziformula The formula specifying the zero-inflation model

#### Value

Returns a list.

```
# Use vitamind data but add a "patient" identifier used as a random effect
vitamind2 <- vitamind</pre>
vitamind2$patient <- sample(</pre>
  LETTERS[1:5],
  size = nrow(vitamind2), replace = TRUE
# Use update_formula_and_data() to perform wrangling steps of cglmm()
# without yet fitting the model
data_and_formula <- update_formula_and_data(</pre>
  data = vitamind2,
  formula = vit_d ~ X + amp_acro(time,
    group = "X",
    period = 12
  )
)
# print formula from above
data_and_formula$newformula
# fit model while adding random effect to cosinor model formula.
mod <- fit_model_and_process(</pre>
  obj = data_and_formula,
  formula = update.formula(
    data_and_formula$newformula, . ~ . + (1 | patient)
  )
)
mod$fit # printing the `glmmTMB` model within shows Std.Dev. of random effect
```

26 vitamind

vitamind

Vitamin D dataset for cosinor modeling examples.

## Description

Simulated data set to illustrate the cosinor model. The vit\_d column contains the blood vitamin D levels which vary over time (time). The rhythm of the vitamind D fluctuations follows a cosine function and can be modeled with a cosinor model. The X column is a binary covariate representing two groups of patients and is associated with the characteristics of the rhythm. The rhythm has a period of about 12 hours.

## Usage

vitamind

#### **Format**

A data.frame with 3 variables: vit\_d, time, and X.

# **Index**

```
* datasets
    cosinor_mixed, 7
    vitamind, 26
amp\_acro, 2, 6
\verb"autoplot.cglmm", 4"
cglmm, 6
cosinor_mixed, 7
fit_model_and_process, 8
polar_plot, 9
{\tt polar\_plot.cglmm}, \\ 11
predict.cglmm, 14
\verb|print.cglmm|, 14|
print.cglmmSubTest, 15
print.cglmmSummary, 16
print.cglmmTest, 17
\verb|sigma.cglmm|, 18|
simulate_cosinor, 19
\verb|summary.cglmm|, 20|\\
test_cosinor_components, 21
test_cosinor_levels, 23
update\_formula\_and\_data, \\ 24
vitamind, 26
```