Package 'ggetho'

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|---|--|
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| Description Extension of 'ggplot2' providing layers, scales and preprocessing functions useful to represent behavioural variables that are recorded over multiple animals and days. This package is part of the rethomics framework http://rethomics.github.io/ >. | |
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geom_peak

Visualise peaks in a power spectrum or periodogram

Description

This function draws points on the x-y coordinates of selected peaks and write their (y) value on the bottom of the plot.

Usage

```
geom_peak(mapping = NULL, data = NULL, stat = "identity",
   position = "identity", ..., na.rm = TRUE, show.legend = NA,
   inherit.aes = TRUE, peak_rank = 1, conversion = hours)
```

| mapping | Set of aesthetic mappings created by aes() or aes_(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping. |
|----------|---|
| data | The data to be displayed in this layer. There are three options: |
| | If NULL, the default, the data is inherited from the plot data as specified in the call to ggplot(). |
| | A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify() for which variables will be created. |
| | A function will be called with a single argument, the plot data. The return value must be a data.frame., and will be used as the layer data. |
| stat | The statistical transformation to use on the data for this layer, as a string. |
| position | Position adjustment, either as a string, or the result of a call to a position adjustment function. |
| | Other arguments passed on to layer(). These are often aesthetics, used to set an aesthetic to a fixed value, like color = "red" or size = 3. They may also be parameters to the paired geom/stat. |
| na.rm | If FALSE, the default, missing values are removed with a warning. If TRUE, missing values are silently removed. |

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| show.legend | logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display. |
|-------------|--|
| inherit.aes | If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. borders(). |
| peak_rank | numerical vector specifying the rank(s) of peak(s) to draw |
| conversion | function to convert values of x to a specific unit. The default, hours, will write x (time) in decimal hours. |

Details

In the input data, peaks are encoded as an additional column/aesthetic with values corresponding to peak ranks (and 0 when the point is not a peak). In other word, the mapping must provide x, y and peak. Only peaks matching peak_rank will be drawn (see example).

References

• The relevant rethomic tutorial section

See Also

- ggperio to create a periodogram
- zeitgebr::find_peaks to automatically add a peak column on periodogram data

Other layers: stat_bar_tile_etho, stat_ld_annotations, stat_pop_etho

```
# We make a data frame by hand with five rows
# There are two peaks: in position 4 and 2
df \leftarrow data.frame(x = hours(1:5),
                 y = c(1, 2, 0, 4, 1),
                 peak = c(0, 2, 0, 1, 0)
# We draw the plot as a line
pl \leftarrow ggplot(df, aes(x, y, peak = peak)) +
                  geom_line() +
                  scale_x_hours()
pl
# Now we could add the peak values as an extra layer:
# The first peak
pl + geom_peak()
# The first ans second peak
pl + geom_peak(peak_rank = 1:2)
# The second only
pl + geom_peak(peak_rank = 2)
# Just like with other geoms,
# we can change colour, size, alpha, shape, ...:
pl + geom_peak(colour = "red", size = 10, alpha = .5, shape = 20)
```

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```
## In the context of circadian analysis,
# Using the zeitgebr package:
require(zeitgebr)
# We make toy data
metadata <- data.table(id = sprintf("toy_experiment|%02d", 1:40),</pre>
                        region_id = 1:40,
                        condition = c("A", "B"),
                        sex = c("M", "M", "F", "F"))
dt <- toy_activity_data(metadata, seed = 107)</pre>
\# We shift period of the group "A" by 0.01
dt[, t := ifelse(xmv(condition) == "A", t, t * 1.01)]
# We compute a periodogram for each individual
per_dt <- periodogram(moving, dt, FUN = chi_sq_periodogram)</pre>
per_dt <- find_peaks(per_dt)</pre>
out <- ggperio(per_dt, aes(y = power - signif_threshold, colour = condition, peak = peak)) +</pre>
                     stat_pop_etho() +
                     facet_wrap( ~ id, labeller = id_labeller)
out
out + geom_peak(colour="black")
```

ggetho

Prepare a ggplot object to represent behavioural data

Description

This function summarises a variable of interest (y or z axis) in order to subsequently represent it over time (x axis) (using layers provided either by ggplot2 or ggetho).

Usage

```
ggetho(data, mapping, summary_FUN = mean,
  summary_time_window = mins(30), time_wrap = NULL, time_offset = 0,
  multiplot = NULL, multiplot_period = hours(24), ...)
```

Arguments

data

behavr::behavr table containing the data and metadata

mapping

default list of aesthetic mappings to use for plot

summary_FUN

method (function) used to summarise variable over time (typically, the mean)

summary_time_window

width (in seconds) of the time window to compute a summary on

time_wrap

time (in seconds) used to wrap the data (see details)

time_offset

time offset (i.e. phase, in seconds) when using time_wrap

multiplot

integer, greater than two, or NULL, the default (see details)

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```
multiplot_period
the duration of the period when mutiplotting (see details)
... additional arguments to be passed to ggplot2::ggplot()
```

Details

time_wrap is typically used to express time relatively to the start of the day. In other words, it can help be used to pull all days together in one representative day. In this case, time_wrap = hours(24). Instead of representing data from the start of the day, it can be done from any offset, using time_offset. For instance, time_offset = hours(12) puts the circadian reference (ZT0) in the middle of the plot.

Multiplotting is a generalisation of double-plotting, triple-plotting... This type or representation is useful to understand periodic behaviours. When multiplot is *not* NULL, data is repeated as many time as its value, along the x axis. The y axis is then the period (typically the day) onset. It is possible to set duration of the period, which is typically 24 h to arbitrary values using the multiplot_period argument.

Value

An initial plot object that can be further edited.

References

· The relevant rethomic tutorial section

See Also

- stat pop etho to show population trend by aggregating individuals over time
- stat_tile_etho to show variable of interest as colour intensity
- stat_ld_annotations to show light and dark phases on the plot

```
# We start by making a dataset with 20 animals
metadata <- data.table(id = sprintf("toy_experiment|%02d", 1:20),</pre>
                   condition = c("A", "B")
dt <- toy_activity_data(metadata, 3)</pre>
# We build a plot object with **nothing inside** (just the axis)
# we want to show proportion of time sleeping on the y axis:
pl <- ggetho(dt, aes(y = asleep))</pre>
pl
# Sometimes, the variable of interest in not on the y axis, but on z axis (colour scale).
# When we do not provide a y axis,
# ggetho will make an ID fo each animal and display them on separate rows
pl <- ggetho(dt, aes(z = asleep))
pl
# this one is the same type, but it groups the animals by condition
pl <- ggetho(dt, aes(z = asleep, y = condition))</pre>
# sorting with paste
```

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ggperio

Prepare a ggplot object to represent periodogram data

Description

This function summarises periodogram data (containing periodograms of multiple individual), to show period on the x axis, and power (or equivalent) on the y axis.

Usage

```
ggperio(data, mapping = aes(x = period, y = power), ...)
```

Arguments

data behavr::behavr table containing the data and metadata
mapping default list of aesthetic mappings to use for plot
additional arguments to be passed to ggplot2::ggplot()

References

• The relevant rethomic tutorial section

See Also

- ggetho to plot time series
- geom_peak to draw peaks on a periodogram
- zeitgebr::periodogram to compute periodograms in a first place

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Examples

```
require(zeitgebr)
# We make toy data
metadata <- data.table(id = sprintf("toy_experiment|%02d", 1:40),</pre>
                        region_id = 1:40,
                        condition = c("A", "B"),
                        sex = c("M", "M", "F", "F"))
dt <- toy_activity_data(metadata, seed = 107)</pre>
# We shift period of the group "A" by 0.01
dt[, t := ifelse(xmv(condition) == "A", t, t * 1.01)]
# We compute a periodogram for each individual
per_dt <- periodogram(moving, dt, FUN = chi_sq_periodogram)</pre>
# Then we display them as an average
out <- ggperio(per_dt, aes(y = power, colour = condition))</pre>
out + stat_pop_etho()
out <- ggperio(per_dt, aes(y = power - signif_threshold, colour = condition))</pre>
out + stat_pop_etho()
out <- ggperio(per_dt, aes(y = power - signif_threshold, colour = condition))</pre>
out + stat_pop_etho() + facet_wrap( ~ id, labeller = id_labeller)
```

ggspectro

Prepare a ggplot object to represent spectrogram data

Description

This function summarises spectrogram data (containing spectrograms of multiple individual), to show period on the y axis, time on the x axis and power on the z axis (e.g. as a colour).

Usage

```
ggspectro(data, mapping = aes(), summary_FUN = mean,
  summary_time_window = mins(30), time_wrap = NULL, time_offset = 0,
  ...)
```

```
data

behavr::behavr table containing the data and metadata

mapping

default list of aesthetic mappings to use for plot

summary_FUN

method (function) used to summarise variable over time (typically, the mean)

summary_time_window

width (in seconds) of the time window to compute a summary on

time_wrap

time (in seconds) used to wrap the data (see details)

time_offset

time offset (i.e. phase, in seconds) when using time_wrap

additional arguments to be passed to ggplot2::ggplot()
```

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Examples

```
library(zeitgebr)
data(dams_sample)
dt <- dams_sample
spect_dt <- spectrogram(activity, dt)
pl <- ggspectro(spect_dt,time_wrap = hours(24)) + stat_tile_etho() + scale_y_hours(log=T) +
    stat_ld_annotations(ld_colours = c("grey","black"))
pl + facet_grid(period_group ~ .)
pl + facet_wrap(~ id)</pre>
```

id_labeller

A facet labeller for id

Description

This function returns a ggplot2::labeller that displays the id on several lines to improve readability.

Usage

```
id_labeller(labels)
```

Arguments

labels

Data frame of labels. Usually contains only one element, but faceting over multiple factors entails multiple label variables.

See Also

ggplot2::labeller, to make your own labellers

```
library(behavr)
metadata <- data.frame(
    id = sprintf("2017-09-01 20:00:12|toy_experiment_a_very_long_name|%02d", 1:20),
    condition = c("A", "B"))
dt <- toy_activity_data(metadata, duration = hours(2))
pl <- ggetho(dt, aes(y = asleep)) + stat_pop_etho()
## Without labelling
pl + facet_wrap( ~ id)

## With labeller
pl + facet_wrap( ~ id, labeller = id_labeller)</pre>
```

stat_bar_tile_etho 9

| stat_bar_tile_etho | Display a variable of interest either as a colour intensity value or as a bar height |
|--------------------|--|
|--------------------|--|

Description

These functions show the temporal trend (time on the x axis) of a variable of interest (z axis) as either colour intensity (stat_tile_etho) or using the hight of the tiles (stat_bar_tile_etho). In both cases, the y axis is a discrete variable such as a treatment or the id of individuals.

Usage

```
stat_bar_tile_etho(mapping = NULL, data = NULL, geom = "bar_tile",
   position = "identity", ..., method = mean, method.args = list(),
   na.rm = FALSE, show.legend = NA, inherit.aes = TRUE)

stat_tile_etho(mapping = NULL, data = NULL, geom = "raster",
   position = "identity", ..., method = mean, method.args = list(),
   na.rm = FALSE, show.legend = NA, inherit.aes = TRUE)
```

| mapping | Set of aesthetic mappings created by aes() or aes_(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping. |
|-------------|---|
| data | The data to be displayed in this layer. There are three options: |
| | If NULL, the default, the data is inherited from the plot data as specified in the call to ggplot(). |
| | A data. frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify() for which variables will be created. |
| | A function will be called with a single argument, the plot data. The return value must be a data.frame., and will be used as the layer data. |
| geom | The geometric object to use display the data |
| position | Position adjustment, either as a string, or the result of a call to a position adjustment function. |
| | Other arguments passed on to layer(). These are often aesthetics, used to set an aesthetic to a fixed value, like color = "red" or size = 3. They may also be parameters to the paired geom/stat. |
| method | function used to compute the aggregate, when/if grouping several individuals on the same row. The default is fucntion is mean. median, min, max are examples of alternatives. |
| method.args | List of additional arguments passed on to the modelling function defined by method. |

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| na.rm | If FALSE, the default, missing values are removed with a warning. If TRUE, missing values are silently removed. |
|-------------|--|
| show.legend | logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display. |
| inherit.aes | If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. borders(). |

References

• The relevant rethomic tutorial section

See Also

- ggetho to generate a plot object
- stat_pop_etho to show population trend by aggregating individuals over time
- stat_ld_annotations to show light and dark phases on the plot

Other layers: geom_peak, stat_ld_annotations, stat_pop_etho

```
# We start by making a to dataset with 20 animals
metadata <- data.frame(id = sprintf("toy_experiment | %02d", 1:20),</pre>
                   age = c(1, 5, 10, 20),
                   condition = c("A", "B")
print(metadata)
dt <- toy_activity_data(metadata, 3)</pre>
# We build a plot object
pl <- ggetho(dt, aes(z = asleep))</pre>
# A standard plot one row per animal:
pl + stat_tile_etho()
# We can also group animals per condition and calculate the average sleep
pl <- ggetho(dt, aes(z = asleep, y = condition))</pre>
pl + stat_tile_etho()
# We can sort by adding condition AND id on the y axis:
pl <- ggetho(dt, aes(z = asleep, y = interaction(id, condition)))</pre>
pl + stat_tile_etho()
# Same if we want to sort by age
pl <- ggetho(dt, aes(z = asleep, y = interaction(id, age)))</pre>
pl + stat_tile_etho()
# Instead, of the average, maybe we want to show the highest (max)
# posible value of sleep for any time point
pl + stat_tile_etho(method = max)
# We can also use stat_bar_tile as an alternative
pl + stat_bar_tile_etho()
```

stat_ld_annotations 11

Description

This function is used to show light and dark (L and D) phases as boxes on top a plot.

Usage

```
stat_ld_annotations(mapping = NULL, data = NULL,
  position = "identity", ld_colours = c("white", "black"),
  ypos = "bottom", height = 0.03, period = hours(24), phase = 0,
  l_duration = hours(12), outline = "black", x_limits = c(NA, NA),
  ..., na.rm = FALSE, show.legend = FALSE, inherit.aes = TRUE)
```

| 1 6 | iguments | | |
|-----|------------------|--|--|
| | mapping | Set of aesthetic mappings created by <pre>aes()</pre> or <pre>aes_()</pre> . If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping. | |
| | data | The data to be displayed in this layer. There are three options: | |
| | | If NULL, the default, the data is inherited from the plot data as specified in the call to ggplot(). | |
| | | A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify() for which variables will be created. | |
| | | A function will be called with a single argument, the plot data. The return value must be a data.frame., and will be used as the layer data. | |
| | position | Position adjustment, either as a string, or the result of a call to a position adjustment function. | |
| | ld_colours | character vector of length two setting the colours for light and dark phases, respectively. The default is c("white", "black"). | |
| | ypos | position and height of the annotation on the y axis. It can be either "top" or "bottom". The default, "bottom" will put the labels below any data. | |
| | height | relative height of the rectangles. The default is 3 percent (0.03). | |
| | period, phase, l | _duration | |
| | | period, phase and duration of the L phase (in seconds) of the LD cycle. | |
| | outline | colour of the border of the rectangles. A value of NA draws no border. | |
| | x_limits | numerical vector of length 2 for the start and end of the annotations (in seconds). The default, c(NA,NA), uses the full range of the plotted data. | |
| | | Other arguments passed on to layer(). These are often aesthetics, used to set an aesthetic to a fixed value, like color = "red" or size = 3. They may also be parameters to the paired geom/stat. | |
| | | | |

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| na.rm | If FALSE, the default, missing values are removed with a warning. If TRUE, missing values are silently removed. |
|-------------|--|
| show.legend | logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display. |
| inherit.aes | If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. borders(). |

References

• The relevant rethomic tutorial section

See Also

• ggetho to generate a plot object

Other layers: geom_peak, stat_bar_tile_etho, stat_pop_etho

Examples

```
library(behavr)
# We start by making a to dataset with 20 animals
metadata <- data.frame(id = sprintf("toy_experiment | %02d", 1:20),</pre>
                   condition = c("A", "B"))
dt <- toy_activity_data(metadata, 3)</pre>
# We build a plot object
pl <- ggetho(dt, aes(y = asleep)) + stat_pop_etho()</pre>
pl + stat_ld_annotations()
# We can also put the annotations in the background:
pl <- ggetho(dt, aes(y = asleep)) +</pre>
                 stat_ld_annotations(outline = NA) +
                 stat_pop_etho()
pl
# Different colours (e.g. DD)
pl + stat_ld_annotations(ld_colour = c("grey", "black"))
# Shorter period
pl + stat_ld_annotations(period = hours(22), phase = hours(3))
# On a tile plot:
pl <- ggetho(dt, aes(z = asleep)) + stat_tile_etho()</pre>
pl + stat_ld_annotations()
```

stat_pop_etho

Compute and display a population aggregate for a variable of interest

Description

This function displays the temporal (time on the x axis) trend of variable of interest, on the y axis as a line with confidence interval as a shaded area.

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Usage

```
stat_pop_etho(mapping = NULL, data = NULL, geom = "smooth",
  position = "identity", ..., method = mean_se, method.args = list(),
  show.legend = NA, inherit.aes = TRUE)
```

Arguments

mapping Set of aesthetic mappings created by aes() or aes_(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping. The data to be displayed in this layer. There are three options: data If NULL, the default, the data is inherited from the plot data as specified in the call to ggplot(). A data. frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify() for which variables will be created. A function will be called with a single argument, the plot data. The return value must be a data.frame., and will be used as the layer data. geom The geometric object to use display the data Position adjustment, either as a string, or the result of a call to a position adjustposition ment function. Other arguments passed on to layer(). These are often aesthetics, used to set an aesthetic to a fixed value, like color = "red" or size = 3. They may also be parameters to the paired geom/stat. method function used to compute the aggregate and confidence intervals. It should return (y, ymin and ymax). The default is ggplot2::mean_se, which computes the mean + or - standard error. ggplot2::mean_cl_boot can be used instead to generate bootstrap confidence interval instead. List of additional arguments passed on to the modelling function defined by method.args method. show.legend logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display. inherit.aes If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. borders().

References

• The relevant rethomic tutorial section

See Also

- ggetho to generate a plot object
- stat_tile_etho to show variable of interest as colour intensity

- stat_ld_annotations to show light and dark phases on the plot
- ggplot2::stat_smooth to understand how to change the type of confidence interval, line colour and so forth

Other layers: geom_peak, stat_bar_tile_etho, stat_ld_annotations

Examples

```
library(behavr)
metadata <- data.frame(id = sprintf("toy_experiment | %02d", 1:20),</pre>
                   age=c(1, 5, 10, 20),
                   condition=c("A", "B"))
dt <- toy_activity_data(metadata, 3)</pre>
# We build a plot object
pl <- ggetho(dt, aes(y = asleep))</pre>
# A standard plot of the whole population:
pl + stat_pop_etho()
# We can also split by condition, and display the two population on different facets:
pl + stat_pop_etho() + facet_grid(condition ~ .)
# Instead, we can use different colour for separate conditions:
pl <- ggetho(dt, aes(y = asleep, colour = condition))</pre>
pl + stat_pop_etho()
# Sometimes, we also have numeric condition (e.g. age)
pl <- ggetho(dt, aes(y = asleep, colour = age))</pre>
pl + stat_pop_etho()
# We could want to aggreate several days of data to one circadian day (i.e. time wrapping)
# here, we also plot the invert of moving (!moving)
pl <- ggetho(dt, aes(y = !moving), time_wrap = hours(24))</pre>
pl + stat_pop_etho()
```

time_scales

Scales for durations

Description

A set of scales used to represent experimental durations.

Usage

```
scale_x_days(name = "Time", breaks = waiver(),
  minor_breaks = waiver(), labels = waiver(), limits = NULL,
  expand = waiver(), oob = scales::censor, na.value = NA_real_,
  position = "bottom", time_wrap = NULL, unit = "day", log = FALSE)

scale_y_days(name = "Time", breaks = waiver(),
  minor_breaks = waiver(), labels = waiver(), limits = NULL,
  expand = waiver(), oob = scales::censor, na.value = NA_real_,
```

```
position = "left", time_wrap = NULL, unit = "day", log = FALSE)
scale_x_hours(name = "Time", breaks = waiver(),
 minor_breaks = waiver(), labels = waiver(), limits = NULL,
 expand = waiver(), oob = scales::censor, na.value = NA_real_,
 position = "bottom", time_wrap = NULL, unit = "h", log = FALSE)
scale_y_hours(name = "Time", breaks = waiver(),
 minor_breaks = waiver(), labels = waiver(), limits = NULL,
 expand = waiver(), oob = scales::censor, na.value = NA_real_,
 position = "left", time_wrap = NULL, unit = "h", log = FALSE)
scale_x_seconds(name = "Time", breaks = waiver(),
 minor_breaks = waiver(), labels = waiver(), limits = NULL,
 expand = waiver(), oob = scales::censor, na.value = NA_real_,
 position = "bottom", time_wrap = NULL, unit = "s", log = FALSE)
scale_y_seconds(name = "Time", breaks = waiver(),
 minor_breaks = waiver(), labels = waiver(), limits = NULL,
 expand = waiver(), oob = scales::censor, na.value = NA_real_,
 position = "left", time_wrap = NULL, unit = "s", log = FALSE)
```

Arguments

name

The name of the scale. Used as axis or legend title. If waiver(), the default, the name of the scale is taken from the first mapping used for that aesthetic. If NULL, the legend title will be omitted.

breaks One of:

- NULL for no breaks
- waiver() for the breaks specified by date_breaks
- A Date/POSIXct vector giving positions of breaks
- A function that takes the limits as input and returns breaks as output

minor_breaks One of:

- · NULL for no breaks
- waiver() for the breaks specified by date_minor_breaks
- A Date/POSIXct vector giving positions of minor breaks
- A function that takes the limits as input and returns minor breaks as output

labels One of:

- NULL for no labels
- waiver() for the default labels computed by the transformation object
- A character vector giving labels (must be same length as breaks)
- A function that takes the breaks as input and returns labels as output

limits A numeric vector of length two providing limits of the scale. Use NA to refer to the existing minimum or maximum.

| expand | Vector of range expansion constants used to add some padding around the data, to ensure that they are placed some distance away from the axes. Use the convenience function ggplot2::expand_scale() to generate the values for the expand argument. The defaults are to expand the scale by 5% on each side for continuous variables, and by 0.6 units on each side for discrete variables. |
|-----------|---|
| oob | Function that handles limits outside of the scale limits (out of bounds). The default replaces out of bounds values with NA. |
| na.value | Missing values will be replaced with this value. |
| position | The position of the axis. "left" or "right" for vertical scales, "top" or "bottom" for horizontal scales |
| time_wrap | duration (in seconds) used to wrap the labels of the time axis |
| unit | the name of unit (string) to be used in the label (e.g. one could use "second" instead of "s") |
| log | logical, whether axis should be on a log-transformed |

Details

time_wrap is useful, for instance, to express time within a day (ZT), instead of absolute time.

References

• The relevant rethomic tutorial section

See Also

- ggetho to generate a plot object
- ggplot2::scale_x_continuous, the defaut ggplot scale, to understand limits, breaks, labels and name

```
# We generate some data
metadata <- data.frame(id = sprintf("toy_experiment | %02d", 1:20),</pre>
                   condition = c("A","B"))
dt <- toy_activity_data(metadata, 3)</pre>
# Then, a simple plot
pl <- ggetho(dt, aes(y = asleep)) + stat_pop_etho()</pre>
pl + scale_x_hours(breaks = days(c(1, 2)))
pl + scale_x_hours()
pl + scale_x_days(breaks = days(c(1, 2)))
pl + scale_x_days()
# To express time modulus `time_wrap`
# e.g. time n the day
pl + scale_x_hours(time_wrap = hours(24)) +
     coord_cartesian(xlim=c(0, days(2)))
# On a shorter time scale
pl <- ggetho(dt[t < hours(5)], aes(z = asleep)) + stat_tile_etho()</pre>
```

```
pl + scale_x_hours()
pl + scale_x_hours(breaks = hours(1:4))
pl + scale_x_seconds(breaks = hours(1:4))
```

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