# Package 'sleepr'

### August 1, 2018

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|--|
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| <b>Description</b> Use behavioural variables to score activity and infer sleep from bouts of immobility. |
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| R topics documented:   |
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bout\_analysis

bout\_analysis

Find "bouts" in categorical time series

#### **Description**

This function is used to find contiguous regions of unique value in a – potentially irregular/heterogeneous – univariate categorical time series.

#### Usage

```
bout_analysis(var, data)
```

#### Arguments

var name of the variable to use from data

data data.table containing behavioural variable from or one multiple animals. When

it has a key, unique values, are assumed to represent unique inviduals (e.g. in a behavr table). Otherwise, it analysis the data as comming from a single animal.

data must have a column t representing time.

#### Value

an object of the same type as data (i.e. data.table::data.table or behavr::behavr). Each row is a specific bout characterised by three columns.

- t its *onset*
- duration its length
- <var> a column with the same name as var. The value of var for this bout.

#### References

• The relevant rethomic tutorial section – on sleep analysis

#### See Also

- sleep\_annotation to generate a binary sleep variable
- rle run length encoding function on which this analysis is based

#### **Examples**

```
# Bout analysis on binary variable:
dt <- toy_dam_data()
dt[, moving := activity > 0]
bdt <- bout_analysis(moving,dt)
print(bdt)
# With multiple states
dt <- toy_ethoscope_data()
# we discretise x position in three states: left, middle and right (1/3 each)</pre>
```

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curate\_dead\_animals

Remove - irrelevant - data after individual have died

#### **Description**

This function detects when individuals have died base on their first (very) long bout of immobility. Following data (which may include spurious artefact of movement) are removed.

#### Usage

```
curate_dead_animals(data, moving_var = moving, time_window = hours(24),
    prop_immobile = 0.01, resolution = 24)
```

#### **Arguments**

| data                     | data.table containing behavioural variable from or one multiple animals. When it has a key, unique values, are assumed to represent unique inviduals (e.g. in a behavr table). Otherwise, it analysis the data as comming from a single animal. data must have a column t representing time. |
|--------------------------|--|
| moving_var               | logical variable in data used to define the moving (alive) state (default is moving)   |
| time_window              | window during which to define death (default is one day)   |
| <pre>prop_immobile</pre> | proportion of immobility that counts as "dead" during time_window (see details)  |
| resolution               | how much scanning windows overlap. Expressed as a factor (see details).  |

#### **Details**

This function scans the time series looking for "death" in the right (future) data, within time\_window. Death is defined as mean(moving\_var) < prop\_immobile within a time window. Moving window start every time\_window/resolution. resolution = 1 is fast but means no overlap. The default would score an animal as dead it does not move more than *one percent of the time* for at least *one day*. All data following "death" are removed.

#### Value

an object of the same type as data (i.e. data.table::data.table or behavr::behavr).

#### References

• The relevant rethomic tutorial section – on high-resolution survival analysis

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#### See Also

• sleep\_annotation – to score movement and sleep

#### **Examples**

```
dt1 <- toy_activity_data()</pre>
#all movement after day 3 is set at 0
dt1[t > days(3), moving := FALSE]
# one artefact of movement is detected at day 3.5
dt1[t == days(3.5), moving := TRUE]
dt2 <- curate_dead_animals(dt1)</pre>
dt3 <- curate_dead_animals(dt1,prop_immobile = 0)</pre>
## Not run:
library(ggplot2)
ggplot(data=dt1[,test:=1],aes(t, as.numeric(moving))) +
  geom_line(data=dt1[,test:=1]) +
  geom_line(data=dt2[, test:=2])+
  geom_line(data=dt3[, test:=3])+
  facet_grid(test ~ .)+
  scale_x_time()
## End(Not run)
```

motion\_detectors

Motion detector for Ethocope data

#### **Description**

Defines whether a *single animal* is moving according to:

#### Usage

```
max_velocity_detector(data, time_window_length,
    velocity_correction_coef = 0.003, masking_duration = 6)
max_velocity_detector_legacy(data, velocity_threshold = 0.006)
virtual_beam_cross_detector(data, time_window_length)
```

#### Arguments

data

data.table::data.table containing behavioural variables of *a single animal* (no id). It must have the columns xy\_dist\_log10x1000(for computing subpixel velocity), x(beam cross), t and has\_interacted (whether a stimulus was delivered).

time\_window\_length

number of seconds to be used by the motion classifier. This corresponds to the sampling period of the output data.

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```
velocity_correction_coef
```

an empirical coefficient to correct velocity with respect to variable framerate.

masking\_duration

number of second during which any movement is ignored (velocity is set to 0) after a stimulus is delivered (aka interaction).

velocity\_threshold

uncorrected velocity above which an animal is classified as 'moving' (for the legacy version).

#### **Details**

- Validated and corrected subpixel velocity (max\_velocity\_detector), the most rigorous
- Uncorrected subpixel velocity (max\_velocity\_detector\_legacy)
- Crossing a virtual beam in the middle of the region of interest (virtual\_beam\_cross\_detector)

max\_velocity\_detector is the default movement classification for real-time ethoscope experiments. It is benchmarked against human-generated ground truth.

These functions are rarely called directly, but typically used is in the context of sleep\_annotation.

#### Value

an object of the same type as data (i.e. data.table::data.table or behavr::behavr) with additional columns:

- moving Logical, TRUE iff. motion was detected.
- beam\_crosses The number of beam crosses (when the animal crosses x = 0.5 that is the midpoint of the region of interest) within the time window
- max\_velocity The maximal velocity within the time window. The resulting data is sampled
  at a period equals to time\_window\_length.

#### See Also

• sleep\_annotation – which requieres a motion detector

sleep\_annotation

Score sleep behaviour from immobility

#### **Description**

This function first uses a motion classifier to decide whether an animal is moving during a given time window. Then, it defines sleep as contiguous immobility for a minimal duration.

#### Usage

```
sleep_annotation(data, time_window_length = 10, min_time_immobile = 300,
   motion_detector_FUN = max_velocity_detector, ...)
sleep_dam_annotation(data, min_time_immobile = 300)
```

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#### **Arguments**

data

data.table containing behavioural variable from or one multiple animals. When it has a key, unique values, are assumed to represent unique inviduals (e.g. in a behavr table). Otherwise, it analysis the data as comming from a single animal. data must have a column t representing time.

time\_window\_length

number of seconds to be used by the motion classifier. This corresponds to the sampling period of the output data.

min\_time\_immobile

Minimal duration (in s) of a sleep bout. Immobility bouts longer or equal to this value are considered as sleep.

motion\_detector\_FUN

function used to classify movement

... extra arguments to be passed to motion\_classifier\_FUN.

#### **Details**

The default time\_window\_length is 300 seconds – it is also known as the "5 minute rule". sleep\_annotation is typically used for ethoscope data, whilst sleep\_dam\_annotation only works on DAM2 data. These functions are *rarely used directly*, but rather passed as an argument to a data loading function, so that analysis can be performed on the go.

#### Value

a behavr table similar to data with additional variables/annotations (i.e. moving and asleep). The resulting data will only have one data point every time\_window\_length seconds.

#### References

• The relevant rethomic tutorial section – on sleep analysis

#### See Also

- motion detectors options for the motion\_detector\_FUN argument
- bout\_analysis to further analyse sleep bouts in terms of onset and length

#### **Examples**

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```
###### Ethoscope, virutal beam cross classification ########
sleep_dt2 <- sleep_annotation(dt_one_animal,</pre>
                             motion_detector_FUN=virtual_beam_cross_detector)
## Not run:
library(ggplot2)
ggplot(sleep_dt2, aes(t,y="Animal 1",fill=asleep)) +
                                    geom_tile() + scale_x_time()
## End(Not run)
###### DAM data, de facto beam cross classification ######
dt_one_animal <- toy_dam_data(seed=7)</pre>
sleep_dt <- sleep_dam_annotation(dt_one_animal)</pre>
## Not run:
library(ggplot2)
ggplot(sleep_dt, aes(t,y="Animal 1",fill=asleep)) +
                                    geom_tile() + scale_x_time()
## End(Not run)
```

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