Package ‘mousetrap’

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Description Mouse-tracking, the analysis of mouse movements in computerized experiments, is a method that is becoming increasingly popular in the cognitive sciences. The mousetrap package offers functions for importing, preprocessing, analyzing, aggregating, and visualizing mouse-tracking data. An introduction into mouse-tracking analyses using mousetrap can be found in Wulff, Kieslich, Henninger, Haslbeck, & Schulte-Mecklenbeck (2021) <doi:10.31234/osf.io/v685r> (preprint: <https://psyarxiv.com/v685r>).

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Create Bezier-curves using the Bernstein approximation.

Usage

bezier(x = c(0, 1, -1), y = c(0, 1.5, 1.5), w = 1, resol = 100)

Arguments

x a numeric vector giving the x-coordinates of exactly three Bezier-points. Defaults to c(0,1,-1) matching the 'mt' format in mt_align.

y a numeric vector giving the x-coordinates of exactly three Bezier-points. Defaults to c(0,1.5,1.5) matching the 'mt' format in mt_align.

w a numeric value or vector specifying one or several Bezier curves, with w governing the pull towards the middle point. Each entry in w creates one Bezier-curve.

resol a numeric value specifying the spatial resolution of the bezier curves. For example, resol = 100 creates bezier curves comprised of 100 points each.

Value

A trajectory array containing the bezier curves.

Author(s)

Dirk U. Wulff
Examples

```r
# Generate range of Bezier-curves
bezier_curves <- bezier(w=seq(0,10,.1))

# Plot curves
mt_plot(bezier_curves)
```

`bimodality_coefficient`

Calculate bimodality coefficient.

Description

Calculate the bimodality coefficient for a numeric vector as specified in Pfister et al. (2013).

Usage

`bimodality_coefficient(x, na.rm = FALSE)`

Arguments

- `x`: a numeric vector.
- `na.rm`: logical specifying whether missing values should be removed.

Details

The calculation of the bimodality coefficient involves calculating the skewness and kurtosis of the distribution first. For this, the `skew` and `kurtosi` functions of the psych package are used. Note that type is set to "2" for these functions in accordance with Pfister et al. (2013).

Value

A numeric value.

Author(s)

Pascal J. Kieslich
Felix Henninger

References

See Also

- *skew* for calculating skewness and kurtosis.
- *mt_check_bimodality* for assessing bimodality using several methods in a mousetrap data object.

Examples

```r
pfister_data_a <- rep(1:11, times=c(3,5,5,10,17,20,17,10,5,5,3))
bimodality_coefficient(pfister_data_a) # .34
pfister_data_b <- rep(1:11, times=c(2,26,14,6,2,0,2,6,14,26,2))
bimodality_coefficient(pfister_data_b) # .79
```

### Description

A data object of class "mousetrap" with the imported and preprocessed mouse-tracking data from Kieslich & Henninger (2017). More information about the study and raw data can be found in `KH2017_raw`.

### Usage

`KH2017`

### Format

A mousetrap data object is a list containing at least the following objects:

- **data**: a `data.frame` containing the trial data (from which the mouse-tracking data columns have been removed). More information about the content of the trial data in KH2017 can be found in `KH2017_raw`. The *rownames* of `data` correspond to the trial identifier. For convenience, the trial identifier is also stored in an additional column called "mt_id".

- **trajectories**: an `array` containing the raw mouse-tracking trajectories. The first dimension represents the different trials and the dimension names (which can be accessed using *rownames*) correspond to the trial identifier (the same identifier that is used as the *rownames* in `data`). The second dimension corresponds to the samples taken over time which are included in chronological order. The third dimension corresponds to the different mouse-tracking variables (timestamps, x-positions, y-positions) which are usually called *timestamps*, *xpos*, and *ypos*.

Some functions in this package (e.g., `mt_time_normalize` and `mt_average`) add additional trajectory arrays (e.g., `tn_trajectories` and `av_trajectories`) to the mousetrap data object. Other functions modify the existing arrays (e.g., `mt_derivatives` adds distance, velocity, and acceleration to an existing dataset). Finally `mt_measures` adds an additional data.frame with mouse-tracking measures to it.
Details

The raw dataset (KH2017_raw) was filtered keeping only correctly answered trials. The filtered dataset was imported using `mt_import_mousetrap`. Trajectories were then remapped using `mt_remap_symmetric` so that all trajectories end in the top-left corner and their starting point was aligned to a common value (0,0) using `mt_align_start`.

References


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**KH2017_raw**

Raw mouse-tracking dataset from Kieslich & Henninger (2017)

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Description

Raw mouse-tracking dataset from Kieslich & Henninger (2017), an experiment using the material and procedure of experiment 1 by Dale et al. (2007). A preprocessed (as opposed to raw) version of the same data can be found in KH2017.

Usage

KH2017_raw

Format

A data.frame with 1140 rows and 19 variables. The data.frame is based on the combined raw data that were created using `read_opensesame` from the `readbulk` library. For ease of use, unnecessary columns were excluded.

The variables included relate to the item that was presented (Exemplar), the answer categories (Category1 and Category2), the subject identifier (subject_nr), the subjects’ response (response), the correctness of the response (response) as well as the mouse-tracking variables (timestamps_get_response, xpos_get_response and ypos_get_response)

Each mouse-tracking variable contains a list of values (separated by ‘,’ ‘)’

- one entry for each recorded position of the mouse. The position coordinates are given in pixels, such that values of zero for both xpos_get_response and ypos_get_response indicate that the cursor is located in the center of the screen. Both variables increase in value as the mouse moves toward the bottom right. Timestamps are given in milliseconds.
Details

The data stem from a study by Kieslich & Henninger (2017) which used the material and procedure of experiment 1 by Dale et al. (2007). In this experiment, participants have to assign exemplars (e.g., "whale") to one of two categories (e.g., "fish" or "mammal") by clicking on the button corresponding to the correct category. All exemplars and categories from the original study were translated to and presented in German.

The data was collected in OpenSesame using the mousetrap plugin (Kieslich & Henninger, 2017). Across the 19 trials of the experiment, 60 participants categorized 13 exemplars that were typical of their category and 6 atypical exemplars for which this was not the case. For the atypical exemplars (e.g., "whale"), the competing category ("fish") was selected to compete with the correct category ("mammal"). The hypothesis under investigation is whether participants’ mouse trajectories deviate more towards the competing category for the atypical exemplars, indicating increased conflict between the response options.

References


Description

The mousetrap package provides functions for importing, preprocessing, analyzing, aggregating, and visualizing mouse-tracking data. In the following, a brief overview of the functions in this package is given.

Read functions

Depending on the file format, one of the standard R functions for reading files into R can be used (e.g., read.table or read.csv).

If raw data were collected using MouseTracker, the mousetrap package provides the read_mt function to read files in the ".mt" format.

If several raw data files should be read and merged, the read_bulk function from the readbulk package can be used (or the read_opensesame function, if data were collected using OpenSesame).

Import functions

The initial step to prepare data for analysis in the mousetrap package is to create a mousetrap data object. Depending on the input format, one of the following functions can be used. A detailed description (and example) of the resulting mousetrap data object can be found in mt_example.
mt_import_mousetrap imports mouse-tracking data that were recorded using the mousetrap plugin for OpenSesame.

mt_import_wide imports mouse-tracking data saved in a wide format (e.g., data collected using MouseTracker).

mt_import_long imports mouse-tracking data saved in a long format. (e.g., trajectories exported using mt_export_long).

**Geometric preprocessing functions**

A number of functions are available that perform geometric preprocessing operations.

mt_remap_symmetric remaps mouse trajectories to one side (or one quadrant) of the coordinate system.

mt_align is a general purpose function for aligning and rescaling trajectories. For specific operations, you can rely on one of the following functions.

mt_align_start aligns the start position of trajectories.

mt_align_start_end aligns all trajectories so that they share a common initial and final coordinate (this is also sometimes referred to as "space-normalization").

**Resampling and interpolation functions**

A number of functions are available that perform resampling and interpolation operations.

mt_exclude_initiation excludes the initial phase of a trial without mouse movement.

mt_exclude_finish excludes a potential phase without mouse movement at the end of a trial.

mt_time_normalize performs time-normalization using equidistant time intervals, resulting in an identical number of samples for all trajectories.

mt_resample resamples trajectories so that samples occur at constant intervals of a specified length.

mt_average averages trajectory coordinates (and related variables) for time bins of constant duration.

mt_length_normalize re-represents each trajectory spatially so that adjacent points on the trajectory become equidistant to each other.

**Data handling functions**

A number of functions are available for data handling operations, such as filtering or adding of new variables or trajectories.

mt_subset filters mouse-tracking data by trials, so that only those meeting the defined criteria are included.

mt_add_variables adds new, self created variables to a trajectory array.

mt_add_trajectory adds a new trajectory to a trajectory array.

mt_bind joins two trajectory arrays.
Analysis functions

A number of different analysis procedures and summary statistics for mouse trajectories have been established in the existing literature. The following functions implement many of these approaches.

- `mt_derivatives` calculates distance, velocity, and acceleration for trajectories.
- `mt_angles` calculates movement angles for trajectories.
- `mt_deviations` calculates the deviations from an idealized trajectory (straight line).
- `mt_measures` calculates a set of mouse-tracking measures.
- `mt_sample_entropy` calculates sample entropy.
- `mt_standardize` standardizes mouse-tracking measures onto a common scale (separately for subsets of the data, e.g., per participant).
- `mt_scale_trajectories` provides different options for standardizing variables in a mouse trajectory array.
- `mt_check_bimodality` assesses the bimodality of mouse-tracking measure distributions.
- `mt_check_resolution` checks the (temporal) logging resolution of raw trajectories.
- `mt_count` counts the number of observations for each trajectory.

Cluster functions

A number of different functions for clustering trajectories is provided.

- `mt_distmat` computes the dissimilarity/distance between each pair of trajectories.
- `mt_cluster` performs trajectory clustering with a specified number of clusters.
- `mt_cluster_k` estimates the optimal number of clusters using various methods.
- `mt_map` maps trajectories onto a predefined set of prototype trajectories (a core set is provided in `mt_prototypes`).

Reshaping, aggregation, and export functions

A number of helper functions are provided for aggregating, plotting, and exporting the multi-dimensional mouse trajectory arrays.

- `mt_reshape` is a general purpose reshaping and aggregation function for mousetrap data.
- `mt_aggregate` aggregates mouse-tracking data per condition.
- `mt_aggregate_per_subject` aggregates mouse-tracking data per (within subjects-) condition separately for each subject.
- `mt_export_long` exports mouse-tracking data in long format.
- `mt_export_wide` exports mouse-tracking data in wide format.

Visualization functions

The following functions can be used for plotting trajectory data, e.g., individual and aggregated trajectories or velocity profiles.

- `mt_plot` plots individual trajectory data.
- `mt_plot_aggregate` plots aggregated trajectory data.
mt_plot_add_rect adds rectangles to a trajectory plot.
mt_plot_riverbed plots the relative frequency of a selected variable across time.
mt_plot_per_trajectory creates a pdf with separate plots per trajectory.
mt_heatmap and mt_heatmap_ggplot plot trajectory heatmaps.
mt_diffmap for creating a difference-heatmap of two trajectory heatmap images.
mt_animate creates a gif trajectory animation.

Helper functions

bimodality_coefficient calculates the bimodality coefficient.
scale_within scales and centers variables within the levels of another variable.
bezier creates Bezier-curves using the Bernstein approximation.

Datasets

mt_example and mt_example_raw contain a mouse-tracking example dataset for demonstrations using the mousetrap package.

Examples

```r
## Not run:
KH2017 <- mt_import_mousetrap(subset(KH2017_raw,correct==1))
KH2017 <- mt_remap_symmetric(KH2017)
KH2017 <- mt_align_start(KH2017)

## End(Not run)

KH2017 <- mt_time_normalize(KH2017)
KH2017 <- mt_measures(KH2017)

mt_aggregate(
  KH2017, use="measures",
  use_variables=c("MAD", "AD"),
  use2_variables="Condition",
  subject_id="subject_nr"
)

mt_plot_aggregate(KH2017,
  use="tn_trajectories",
  x="xpos", y="ypos", color="Condition",
  subject_id="subject_nr"
)

## Not run:
mt_plot(KH2017,
  use="tn_trajectories",
  x="xpos", y="ypos", color="Condition"
)
```
## End(Not run)

---

### mt_add_trajectory

*Add new trajectory to trajectory array.*

### Description

Add a single new trajectory to trajectory array.

### Usage

```r
mt_add_trajectory(
  data,
  use = "trajectories",
  save_as = use,
  xpos = NULL,
  ypos = NULL,
  xypos = NULL,
  id = "new"
)
```

### Arguments

- **data**: a mousetrap data object created using one of the `mt_import` functions (see `mt_example` for details). Alternatively, a trajectory array can be provided directly (in this case `use` will be ignored).
- **use**: a character string specifying which trajectory data should be used.
- **save_as**: a character string specifying where the resulting trajectory data should be stored.
- **xpos**: a vector of x positions. Ignored, if `xypos` is provided.
- **ypos**: a vector of y positions. Ignored, if `xypos` is provided.
- **xypos**: a matrix, the first column corresponding to the x positions, the second to the y positions.
- **id**: a character string specifying the identifier of the to be added trajectory.

### Value

A mousetrap data object (see `mt_example`) where the new trajectory has been added. If the trajectory array was provided directly as `data`, only the trajectory array will be returned.

### Author(s)

- Pascal J. Kieslich
- Felix Henninger
**mt_add_variables**

**Description**

Add new variables to the trajectory array (and remove potentially existing variables of the same name). This is mostly a helper function used by other functions in this package (e.g., `mt_deviations`). However, it can also be helpful if the user has calculated new variables for each logged coordinate and wants to add them to an existing trajectory array.

**Usage**

```r
mt_add_variables(data, use = "trajectories", save_as = use, variables)
```

**Arguments**

- **data**: a mousetrap data object created using one of the `mt_import` functions (see `mt_example` for details). Alternatively, a trajectory array can be provided directly (in this case `use` will be ignored).
- **use**: a character string specifying which trajectory data should be used.
- **save_as**: a character string specifying where the resulting trajectory data should be stored.
- **variables**: a list of matrices that each contain the data of one of the to be added variables. In this case, the new variables with their values are added as a new entry in the trajectory arrays third dimension. Alternatively, a character vector specifying the name of the new variables that should be added to the trajectory array. In this case, the new variables are filled with NAs.

**Value**

A mousetrap data object (see `mt_example`) where the new variables have been added to the trajectory array. Depending on the input to `variables`, the values for the added variables are either NAs or their actual values. If columns of the same name already existed, they have been removed. If the trajectory array was provided directly as `data`, only the trajectory array will be returned.

**Author(s)**

Pascal J. Kieslich
Felix Henninger
Examples

# Calculate new (arbitrary) variables for this example
# ... the sum of the x- and y-positions
xy_sum <- mt_example$trajectories[,"xpos"] + mt_example$trajectories[,"ypos"]
# ... the product of the x- and y-positions
xy_prod <- mt_example$trajectories[,"xpos"] * mt_example$trajectories[,"ypos"]

# Add the new variables to the trajectory array
mt_example <- mt_add_variables(mt_example,
    variables=list(xy_sum=xy_sum, xy_prod=xy_prod))

mt_aggregate

Aggregate mouse-tracking data per condition.

Description

mt_aggregate is used for aggregating mouse-tracking measures (or trajectories) per condition. One or several condition variables can be specified using use2_variables. Aggregation will be performed separately for each level of the condition variables. mt_aggregate is a wrapper function for mt_reshape.

Usage

mt_aggregate(
  data,
  use = "measures",
  use_variables = NULL,
  use2 = "data",
  use2_variables = NULL,
  subject_id = NULL,
  trajectories_long = TRUE,
  ...
)

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use a character string specifying which dataset should be aggregated. The corresponding data are selected from data using data[[use]]. Usually, this value corresponds to either "tn_trajectories" or "measures", depending on whether the time-normalized trajectories or derived measures should be aggregated.

use_variables a character vector specifying the mouse-tracking variables to aggregate. If a data.frame with mouse-tracking measures is provided as data, this corresponds to the column names. If a trajectory array is provided, this argument should
specify the labels of respective array dimensions. If unspecified, all variables will be aggregated.

use2  
a character string specifying where the data containing the condition information can be found. Defaults to "data" as data[["data"]][usually contains all non mouse-tracking trial data. Alternatively, a data.frame can be provided directly.

use2_variables  
a character string (or vector) specifying the variables (in data[[use2]]) across which the trajectories / measures will be aggregated. For each combination of levels of the grouping variable(s), aggregation will be performed separately using summarize_at.

subject_id  
an optional character string specifying the column that contains the subject identifier. If specified, aggregation will be performed within subjects first (i.e., within subjects for all available values of the grouping variables specified in use2_variables).

trajectories_long  
logical indicating if the reshaped trajectories should be returned in long or wide format. If TRUE, every recorded position in a trajectory is placed in another row (whereby the order of the positions is logged in the variable mt_seq). If FALSE, every trajectory is saved in wide format and the respective positions are indexed by adding an integer to the corresponding label (e.g., xpos_1, xpos_2, ...). Only relevant if data[[use]] contains trajectories.

...  
additional arguments passed on to mt_reshape (such as subset).

Value

A data.frame containing the aggregated data.

Author(s)

Pascal J. Kieslich
Felix Henninger

See Also

mt_aggregate_per_subject for aggregating mouse-tracking measures and trajectories per subject.

summarize_at for aggregating data using the dplyr package.

Examples

# Time-normalize trajectories
mt_example <- mt_time_normalize(mt_example)

# Aggregate time-normalized trajectories per condition
average_trajectories <- mt_aggregate(mt_example, use="tn_trajectories", use2_variables="Condition"
# Calculate mouse-tracking measures
mt_example <- mt_measures(mt_example)

# Aggregate measures per condition
average_measures <- mt_aggregate(mt_example, 
  use="measures", use_variables=c("MAD", "AD"), 
  use2_variables="Condition"
)

# Aggregate measures per condition
# first within subjects and then across subjects
average_measures <- mt_aggregate(mt_example, 
  use="measures", use_variables=c("MAD", "AD"), 
  use2_variables="Condition", 
  subject_id="subject_nr"
)

---

mt_aggregate_per_subject

Aggregate mouse-tracking data per condition separately for each subject.

Description

mt_aggregate_per_subject can be used for aggregating mouse-tracking measures (or trajectories) per condition separately for each subject. One or more condition variables can be specified using use2_variables. Aggregation will be performed separately for each level of the condition variables. mt_aggregate_per_subject is a wrapper function for mt_reshape.

Usage

mt_aggregate_per_subject(
  data, 
  use = "measures", 
  use_variables = NULL, 
  use2 = "data", 
  use2_variables = NULL, 
  subject_id, 
  trajectories_long = TRUE, 
  ...
)

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).
use a character string specifying which dataset should be aggregated. The corresponding data are selected from data using data[[use]]. Usually, this value corresponds to either "tn_trajectories" or "measures", depending on whether the time-normalized trajectories or derived measures should be aggregated.

use_variables a character vector specifying the mouse-tracking variables to aggregate. If a data.frame with mouse-tracking measures is provided as data, this corresponds to the column names. If a trajectory array is provided, this argument should specify the labels of respective array dimensions. If unspecified, all variables will be aggregated.

use2 a character string specifying where the data containing the condition information can be found. Defaults to "data" as data[["data"]]] usually contains all non mouse-tracking trial data. Alternatively, a data.frame can be provided directly.

use2_variables a character string (or vector) specifying the variables (in data[[use2]]) across which the trajectories / measures will be aggregated. For each combination of levels of the grouping variable(s), aggregation will be performed separately using summarize_at.

subject_id a character string specifying which column contains the subject identifier.

trajectories_long logical indicating if the reshaped trajectories should be returned in long or wide format. If TRUE, every recorded position in a trajectory is placed in another row (whereby the order of the positions is logged in the variable mt_seq). If FALSE, every trajectory is saved in wide format and the respective positions are indexed by adding an integer to the corresponding label (e.g., xpos_1, xpos_2, ...). Only relevant if data[[use]] contains trajectories.

Value
A data.frame containing the aggregated data.

Author(s)
Pascal J. Kieslich
Felix Henninger

See Also
mt_aggregate for aggregating mouse-tracking measures and trajectories per condition.
summarize_at for aggregating data using the dplyr package.

Examples
# Time-normalize trajectories
mt_example <- mt_time_normalize(mt_example)

# Aggregate time-normalized trajectories per condition
# separately per subject
```r
average_trajectories <- mt_aggregate_per_subject(
  mt_example,
  use="tn_trajectories",
  use2_variables="Condition",
  subject_id="subject_nr"
)

# Calculate mouse-tracking measures
mt_example <- mt_measures(mt_example)

# Aggregate measures per condition
# separately per subject
average_measures <- mt_aggregate_per_subject(
  mt_example,
  use="measures",
  use_variables=c("MAD", "AD"),
  use2_variables="Condition",
  subject_id="subject_nr"
)
```

---

**mt_align**

**Align trajectories.**

**Description**

*mt_align* aligns trajectories to a common start point, end point, and / or coordinate system.

**Usage**

```r
mt_align(
  data,
  use = "trajectories",
  save_as = use,
  dimensions = c("xpos", "ypos"),
  coordinates = "isotropic",
  align_start = FALSE,
  align_end = FALSE,
  align_side = "no",
  verbose = FALSE
)
```

**Arguments**

- **data**
  - a mousetrap data object created using one of the *mt_import* functions (see *mt_example* for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

- **use**
  - a character string specifying which trajectory data should be used.
mt_align

save_as a character string specifying where the resulting trajectory data should be stored.
dimensions a character string specifying which trajectory variables should be used. Can be of length 2 or 3 for two-dimensional or three-dimensional alignment respectively.
coordinates either a numeric vector of length 4 specifying the xstart, ystart, xend, yend coordinates of the trajectory start and end points. Can also be isotropic (the default) to preserve the coordinates of dim1 and dim2. isotropic-norm to set the coordinates to c(0,0,-1,x) where x is chosen to preserve the aspect ratio of dim1 and dim2. mt to set coordinates to c(0,0,-1,1.5). norm to set coordinates to c(0,0,-1,1) and wide to set coordinates to c(0,0,-1,1.2). In the three-dimensional case, coordinates is a vector of length 6.
align_start boolean specifying whether the start points of all trajectories should be aligned to the position specified in coordinates. See Details.
align_end boolean specifying whether the end points of all trajectories should be aligned to the position specified in coordinates. See Details.
align_side character string specifying whether all trajectories should be flipped to the left side (left), the right side (right), or not at all (no). Assumes that first entry in dimensions are the x positions.
verbose logical indicating whether function should report its progress.

Details

If align_start / align_end is FALSE, coordinates define the position of the average start / end point across all trajectories.

Note that if the end points of trajectories are not aligned, coordinates refer to the hypothetical case where all trajectories are mapped to one side.

If align_start / align_end is TRUE, the start / end point of each trajectory is set to the exact position specified in coordinates. align_start and align_end can be set completely independently of one another, i.e., one can align only end points, only start points, none, or both.

If align_start is set to "left" or "right" trajectories will be flipped to the lower or upper spectrum of the first dimensions, respectively. If the first dimension is the x-coordinate this is equivalent to flipping the trajectories to the left and right side, respectively.

Value

A mousetrap data object (see mt_example) with aligned trajectories. Per default, the dimensions in the original trajectory array will be replaced. If a different trajectory array is specified using save_as, a new trajectory array will be created (including only the aligned dimensions). If a trajectory array was provided directly as data, only the aligned trajectories will be returned.

Author(s)

Dirk U. Wulff
See Also

- `mt_align_start` for aligning all trajectories to a common start position.
- `mt_align_start_end` for aligning all trajectories so that they share a common initial and final coordinate.
- `mt_remap_symmetric` for remapping trajectories to one side (or one quadrant) of the coordinate system.

Examples

```r
mt_example <- mt_align(mt_example, 
                        align_start = TRUE, align_end = TRUE, 
                        coordinates = 'mt')
```

---

### mt_align_start

**Align start position of trajectories.**

Adjust trajectories so that all trajectories have the same start position.

#### Usage

```r
mt_align_start( 
  data, 
  use = "trajectories", 
  save_as = use, 
  dimensions = c("xpos", "ypos"), 
  start = c(0, 0), 
  verbose = FALSE
)
```

#### Arguments

- **data**: a mousetrap data object created using one of the mt_import functions (see `mt_example` for details). Alternatively, a trajectory array can be provided directly (in this case `use` will be ignored).
- **use**: a character string specifying which trajectory data should be used.
- **save_as**: a character string specifying where the resulting trajectory data should be stored.
- **dimensions**: a character vector specifying the dimensions in the trajectory array that should be aligned.
- **start**: a numeric vector specifying the start values for each dimension, i.e., the values the first recorded position should have in every trial. If NULL, trajectories are aligned to the mean first position across all trials.
- **verbose**: logical indicating whether function should report its progress.
Value

A mousetrap data object (see mt_example) with aligned trajectories. All other trajectory dimensions not specified in dimensions (e.g., timestamps) will be kept as is in the resulting trajectory array. If the trajectory array was provided directly as data, only the trajectory array will be returned.

Author(s)

Pascal J. Kieslich
Felix Henninger

See Also

mt_align_start_end for aligning the start and end position of trajectories.
mt_align as a general purpose function for aligning and rescaling trajectories.
mt_remap_symmetric for remapping trajectories.

Examples

```r
# Import raw trajectories for demonstration
mt_example <- mt_import_mousetrap(mt_example_raw)

# Align trajectories to start coordinates (0,0)
mt_example <- mt_align_start(mt_example, start=c(0,0))

# Import raw trajectories for demonstration
mt_example <- mt_import_mousetrap(mt_example_raw)

# Align trajectories to mean first coordinates
mt_example <- mt_align_start(mt_example, start=NULL)
```

Description

Adjust trajectories so that all trajectories have an identical start and end point. In some articles, this is also referred to as space-normalization (e.g. Dale et al., 2007).
mt_align_start_end

Usage

mt_align_start_end(
  data,
  use = "trajectories",
  save_as = use,
  dimensions = c("xpos", "ypos"),
  start = c(0, 0),
  end = c(-1, 1),
  verbose = FALSE
)

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use a character string specifying which trajectory data should be used.

save_as a character string specifying where the resulting trajectory data should be stored.

dimensions a character vector specifying the dimensions in the trajectory array that should be aligned.

start a numeric vector specifying the start values for each dimension, i.e., the values the first recorded position should have in every trial. If NULL, trajectories are aligned to the mean first position across all trials.

end a numeric vector specifying the end values for each dimension, i.e., the values the last recorded position should have in every trial. If NULL, trajectories are aligned to the mean last position across all trials. Note that in this case trajectories should be remapped to one side before alignment (e.g., using mt_remap_symmetric) so that the mean last position is meaningful.

verbose logical indicating whether function should report its progress.

Value

A mousetrap data object (see mt_example) with aligned trajectories. All other trajectory dimensions not specified in dimensions (e.g., timestamps) will be kept as is in the resulting trajectory array. If the trajectory array was provided directly as data, only the trajectory array will be returned.

Author(s)

Pascal J. Kieslich
Felix Henninger

References

See Also

mt_align_start for aligning the start position of trajectories.
mt_align as a general purpose function for aligning and rescaling trajectories.
mt_remap_symmetric for remapping trajectories.

Examples

# Align start and end positions to specific coordinates
mt_example <- mt_align_start_end(mt_example,
                                   start=c(0,0), end=c(-1,1))

# Import raw trajectories for demonstration
mt_example <- mt_import_mousetrap(mt_example_raw)

# Remap trajectories
mt_example <- mt_remap_symmetric(mt_example)

# Align trajectories to mean first and last coordinates
mt_example <- mt_align_start_end(mt_example,
                                   start=NULL, end=NULL)

mt_angles

Calculate movement angles.

Description

Calculate point-based and vertical-based angles for the points in the movement trajectory. Point-based angles are the angle defined by three subsequent points on the trajectory. Vertical-based angles are the angles between two subsequent points and the vertical axis.

Usage

mt_angles(
  data,
  use = "trajectories",
  dimensions = c("xpos", "ypos"),
  save_as = use,
  na_replace = FALSE,
  unit = "radian",
  verbose = FALSE
)

Arguments

- **data**: a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).
- **use**: a character string specifying which trajectory data should be used.
- **dimensions**: a character string specifying which trajectory variables should be used. Must be of length 2.
- **save_as**: a character string specifying where the resulting trajectory data should be stored.
- **na_replace**: logical specifying whether NAs in the angle values should be replaced using the next existing angle value (see Details). Defaults to FALSE.
- **unit**: character specifying the unit for the angles. Default is "radian", alternative is "degree".
- **verbose**: logical indicating whether function should report its progress.

Details

By default, angles are reported in radians, the alternative is degrees. For the first point in a trajectory, the angle values are always not defined (NA).

For vertical-based angles (angle_v), positive values indicate a movement to the left of the vertical, negative values to the right of the vertical. If there was no movement across two consecutive points, angle_v is not defined and, by default, NA is returned. If na_replace is TRUE, the next existing angle value is reported instead.

For point-based angles (angle_p), angles indicate changes of movement within three consecutive time steps. The reported angle is always the smaller one. A value of pi (= 3.14...) (for radians) or 180 (for degrees) indicates a constant movement direction, a value of 0 (both for radians and degrees) a complete reversal. If there was no movement across two consecutive points, angle_p is not defined and, by default, NA is returned. If na_replace is TRUE, the next existing angle value is reported instead. angle_p is also not defined for the last point of the trajectory.

Value

A mousetrap data object (see mt_example) with point-based and vertical-based angles added as additional variables to the trajectory array (called angle_p and angle_v). If a trajectory array was provided directly as data, only the trajectory array will be returned.

Author(s)

Dirk U. Wulff

Examples

```r
# Calculate movement angles
mt_example <- mt_angles(mt_example)

# Calculate movement angles (in degree) and replace NAs with next existing value
mt_example <- mt_angles(mt_example,
```
mt_animate

Create gif trajectory animation.

Description

[Experimental]

mt_animate animates trajectories using the animation package. Note that this function has beta status.

Usage

mt_animate(
  data,
  use = "trajectories",
  dimensions = c("xpos", "ypos"),
  timestamps = "timestamps",
  filename = "trajectory_animation.gif",
  xres = 1000,
  seconds = 3,
  framerate = 24,
  speed = 0.5,
  density = 3,
  jitter = TRUE,
  remove = FALSE,
  bg = "black",
  col = "white",
  lwd = 1,
  loop = FALSE,
  bounds = NULL,
  norm = FALSE,
  upscale = 1,
  decay = 10,
  max_intensity = 5,
  discard_images = TRUE,
  im_path = NULL,
  parallel = TRUE,
  verbose = FALSE
)

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).
use  a character string specifying which trajectory data should be used.
dimensions  a character vector specifying the two dimensions in the trajectory array that contain the mouse positions. Usually (and by default), the first value in the vector corresponds to the x-positions (xpos) and the second to the y-positions (ypos).
timestamps  a character string specifying the trajectory dimension containing the timestamps. If NULL linearly increasing timestamps are assumed, producing a perfectly constant timestamp interval.
filename  character string specifying the path and filename of the resulting .gif. If the extension of filename is not .gif, .gif is added at the end. Must not contain spaces.
xres  numeric specifying the resolution of the .gif file.
seconds  numeric specifying the duration of the .gif file.
framerate  numeric specifying the framerate of the .gif file. Defaults to 24 implying smooth non-discrete frame transitions for the human eye.
speed  numeric specifying the speed of the trajectories with regard to their original velocity profile. I.e., a value of .5 shows trajectories in half of the original velocities, whereas a value of 2 shows trajectories in double of the original velocities.
density  integer specifying the number of trajectories to be added each frame. I.e., if density = 10, seconds = 10, framerate = 24 and speed = .5 then the animation will show 10 x 10 x 24 x .5 = 1200 trajectories.
jitter  logical specifying whether the density should be jittered. If TRUE, density varies according to rgeom (1/density).
remove  logical specifying whether trajectories that reached their end points should be removed from the rest of the animation. Defaults to FALSE implying that all finished trajectories remain visible.
bg  character string specifying the background color.
col  character string specifying the foreground color, i.e., the color used to draw the trajectories.
lwd  numeric specifying the line width of the trajectories.
loop  logical specifying whether gif should be looped. If FALSE (the default), the last frame will remain visible after the animation is finished. If TRUE, the gif will infinitely repeat itself.
bounds  numeric vector specifying the xleft, xright, ybottom, and ytop limits of the animation canvas. Defaults to NULL in which case the animation canvas is set to include all existing trajectory points, irrespective of how extreme they may be.
norm  logical specifying whether the trajectories should be remapped to the mt-space. See mt_align. Note that aligning often requires that that all trajectories are flipped to one side first (see mt_remap_symmetric).
upscale  numeric specifying a scaling factor for the animation resolution. E.g, upscale = 2 implies that the x-resolution in .gif file is 2*xres.
decay  numeric defining a within-trajectory gradient of color intensity. Specifically, values larger than 1 will give more recent movements higher color intensities than movements that lie longer in the past, and vice versa.
max_intensity numeric specifying the maximum color intensity. A value of, e.g., 5, implies that color intensity is limited to 5 overlapping trajectories. I.e., a point at which 4 trajectories overlap will in that case have a smaller color intensity than a point at which 5 trajectories overlap, but there will be no difference between the latter and a point at which 6 trajectories overlap. If decay is unequal 1, this metric refers to the most intense color point within the trajectory.

discard_images logical specifying whether the temporary folder containing the temporary .png images should be deleted. Defaults to TRUE.

im_path character string specifying the location of ImageMagick’s convert function. If NULL, the convert function is expected in ’/usr/local/bin/convert’, the default location for Linux and OSX operating systems. The location has to be specified explicitly for Windows (see Details and Examples).

parallel logical specifying whether the temporary .png images should be created using parallel processing (uses clusterApplyLB). Process will be run on the maximum number of available cores (as determined by detectCores).

verbose logical indicating whether function should report its progress.

Details

mt_animate produces a .gif file showing a continuous stream of animated trajectories. The function first produces a series of .png images, which then are combined into a .gif animation using ImageMagick (see https://www.imagemagick.org/).

In order to run this function, ImageMagick must be installed (download from https://www.imagemagick.org/). Under Unix systems (Linux and Apple’s OSX) the function will look for ImageMagick using its default installation path. Alternatively, the location of ImageMagick’s convert function can be provided using the im_path argument. Under Windows, im_path must always be specified explicitly (e.g., it might look something like this im_path = ”C:/Program Files/ImageMagick-7.0.5-Q16/convert.exe”).

During the animation trajectories are sampled from the data without replacement. The function stops when it reaches the last trajectory contained in data.

By default, mt_animate animates trajectories using the original timestamps. Timestamps are expected to be expressed in milliseconds. By setting timestamps = NULL, the function can also assume timestamps to be regular, i.e., of constant interval, in this case the longest duration is set to exactly one second.

In order to create high-resolution (large) animations in a relatively short time increase upscale in favor of xres. However, note that this will decrease the sharpness of the image.

In order to increase or decrease the overall color intensity decrease or increase the max_intensity, respectively.

Author(s)

Dirk U. Wulff

Examples

## Not run:
# Preprocess trajectory data
mt_example <- mt_align_start(mt_example)
mt_example <- mt_remap_symmetric(mt_example)

# Create animated trajectory gif
# (under Linux / OSX)
mt_animate(mt_example, filename = "MyMovie.gif")

# Increase duration and density while decreasing speed
mt_animate(mt_example, filename = "MyMovie2.gif",
    seconds = 10, speed = .3, density = 10)

# Create animated trajectory gif
# (under Windows - ImageMagick version specific example)
mt_animate(mt_example, filename = "MyMovie.gif",
    im_path = "C:/Program Files/ImageMagick-7.0.5-Q16/convert.exe")

## End(Not run)

---

**mt_average**

*Average trajectories across intervals.*

**Description**

Average trajectory data across specified intervals (e.g., constant time intervals). For every specified dimension in the trajectory array (by default, every dimension, i.e., x- and y-position, possibly also velocity and acceleration etc.), the mean value for the respective interval is calculated (see Details for information regarding the exact averaging procedure).

**Usage**

```r
mt_average(
    data,
    use = "trajectories",
    save_as = "av_trajectories",
    dimensions = "all",
    av_dimension = "timestamps",
    intervals = NULL,
    interval_size = 100,
    max_interval = NULL,
    verbose = FALSE
)
```

**Arguments**

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>data</td>
<td>a mousetrap data object created using one of the mt_import functions (see <code>mt_example</code> for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).</td>
</tr>
</tbody>
</table>
use a character string specifying which trajectory data should be used.

save_as a character string specifying where the resulting trajectory data should be stored.

dimensions a character vector specifying the dimensions in the trajectory array that should be averaged. By default ("all"), all trajectory dimensions will be averaged.

av_dimension a character string specifying which values should be used for determining the intervals for averaging ("timestamps" by default).

intervals an optional numeric vector. If specified, these values are taken as the borders of the intervals (interval_size and max_interval are ignored).

interval_size an integer specifying the size of the constant dimension interval.

max_interval an integer specifying the upper limit of the last dimension value that should be included (therefore, it should be a multiple of the interval_size). If specified, only values will be used for averaging where the dimension values are smaller than max_interval. If unspecified (the default), all values will be included.

verbose logical indicating whether function should report its progress.

Details

For each interval, it is first determined which of the values lie within the respective interval of the dimension used for averaging (e.g., timestamps). Intervals are left-open, right-closed (e.g., if values are averaged across constant timestamps of 100 ms, a timestamp of 1200 would be included in the interval 1100-1200 while a timestamp of 1300 would be included in the interval 1200-1300). Then, all values for which the corresponding average dimension values lie within the interval are averaged.

In case the last interval is not fully covered (e.g., if the last timestamp has the value 1250), values for the corresponding interval (1200-1300) will be computed based on the average of the values up to the last existing value.

Note that mt_average assumes that the trajectory variables are recorded with a constant sampling rate (i.e., with a constant difference in the timestamps). If the sampling rate varies considerably, mt_resample should be called before averaging to arrive at equally spaced timestamps. The sampling rate can be investigated using mt_check_resolution.

If average velocity and acceleration are of interest, mt_derivatives should be called before averaging.

Value

A mousetrap data object (see mt_example) with an additional array (by default called av_trajectories) that contains the average trajectory data per dimension interval. If a trajectory array was provided directly as data, only the average trajectories will be returned.

For the dimension values used for averaging (specified in av_dimension), the midpoint of the respective interval is reported, which is helpful for plotting the trajectory data later on. However, this value does not necessarily correspond to the empirical mean of the dimension values in the interval.

Author(s)

Pascal J. Kieslich
Felix Henninger
mt_bind

Join two trajectory arrays

Description

Join two trajectory arrays. This function is mainly used internally, but can be helpful in those (relatively rare) occasions where additional processed trajectory data should be added to another trajectory array.

Usage

mt_bind(trajectories1, trajectories2, verbose = FALSE)

Arguments

trajectories1 a trajectory array (see mt_example).
trajectories2 a trajectory array (see mt_example).
verbose logical indicating whether function should report its progress.

Value

A trajectory array.

Author(s)

Pascal J. Kieslich
Felix Henninger
Examples

```r
trajectories_combined <- mt_bind(
  mt_example$trajectories,
  mt_prototypes
)
```

---

**mt_check_bimodality**  
*Assess bimodality of mouse-tracking measure distributions.*

---

**Description**

Assess bimodality of the distribution of mouse-tracking measures using the bimodality coefficient and Hartigan’s dip statistic (see Details). If bimodality should be assessed separately for different conditions, the corresponding variables can be specified under `grouping_variables`.

**Usage**

```r
mt_check_bimodality(
  data,  
  use = "measures",  
  use_variables = NULL,  
  methods = c("BC", "HDS"),  
  B = 2000,  
  grouping_variables = NULL,  
  ...
)
```

**Arguments**

- **data**  
a mousetrap data object created using one of the `mt_import` functions (see `mt_example` for details).

- **use**  
a character string specifying which data should be used. By default, points to the `measures` data.frame created using `mt_measures`.

- **use_variables**  
a vector specifying for which mouse-tracking measures bimodality should be assessed.

- **methods**  
a character string (or vector) specifying which methods should be used for assessing bimodality (see Details).

- **B**  
an integer specifying the number of replicates used in the Monte Carlo test (only relevant if "HDS_sim" is included in `methods`, see Details).

- **grouping_variables**  
a character string (or vector) specifying one or more variables in `data[["data"]].` If specified, bimodality will be assessed separately for each level of the variable. If unspecified (the default), bimodality is checked across all trials.

- **...**  
additional arguments passed on to `mt_reshape` (such as `subset`).
Details

Different methods have been suggested for assessing the bimodality of mouse-tracking measure distributions, each of which has advantages and disadvantages (see Freeman & Dale, 2013).

Hehman et al. (2015) focus on two specific methods (bimodality coefficient and Hartigan’s dip statistic) which are implemented here.

If methods include BC, the bimodality coefficient is calculated using the `bimodality_coefficient` function in this package. According to Freeman and Ambady (2010), a distribution is considered bimodal if \( BC > 0.555 \).

Note that MouseTracker (Freeman & Ambady, 2010) standardizes variables within each subject before computing the BC. This is also possible here using `mt_standardize` (see Examples).

If methods include HDS, Hartigan’s dip statistic is calculated using the `dip.test` function of the `diptest` package. The corresponding p value (computed via linear interpolation) is returned.

If methods include HDS_sim, Hartigan’s dip statistic is calculated using the `dip.test` function with the additional argument `simulate.p.values=TRUE`. In this case, the p value is computed from a Monte Carlo simulation of a uniform distribution with B (default: 2000) replicates.

Value

A list of several data.frames. Each data.frame contains the value returned by the respective method for assessing bimodality (see Details) - separately per condition (specified in the row dimension) and measure (specified in the column dimension).

Author(s)

Pascal J. Kieslich
Felix Henninger

References


See Also

`bimodality_coefficient` for more information about the bimodality coefficient.
`dip.test` for more information about Hartigan’s dip test.

Examples

```r
# Calculate measures
mt_example <- mt_measures(mt_example)
```
# Assess bimodality for untransformed variables
mt_check_bimodality(mt_example,
    use_variables=c("MAD", "AD"))

# Standardize variables per participant
mt_example <- mt_standardize(mt_example,
    use_variables=c("MAD", "AD"), within="subject_nr")

# Assess bimodality for standardized variables
mt_check_bimodality(mt_example,
    use_variables=c("z_MAD", "z_AD"))

# Assess bimodality with simulated p values for HDS
mt_check_bimodality(mt_example,
    use_variables=c("z_MAD", "z_AD"),
    methods=c("BC", "HDS_sim"))

# Assess bimodality per condition
mt_check_bimodality(mt_example,
    use_variables=c("z_MAD", "z_AD"),
    grouping_variables="Condition")

---

**mt_check_resolution**  
*Check logging resolution by looking at timestamp differences.*

**Description**

*mt_check_resolution* computes the timestamp differences as a measure of the logging resolution. It provides various descriptive statistics to check the logging resolution.

**Usage**

```r
mt_check_resolution(
    data,  
    use = "trajectories",  
    timestamps = "timestamps",  
    desired = NULL,  
    digits = NULL  
)
```

**Arguments**

- **data**: a mousetrap data object created using one of the *mt_import* functions (see *mt_example* for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).
- **use**: a character string specifying which trajectory data should be used.
- **timestamps**: a character string specifying the trajectory dimension containing the timestamps.
mt_cluster

desired an optional integer. If specified, additional statistics are computed concerning the (relative) frequencies with which exactly the desired timestamp difference (with tolerance 1e-12) occurred.

digits an optional integer. If specified, timestamps will be rounded before performing any checks. Potentially useful if timestamps are recorded with submillisecond precision.

Details

If mouse-tracking experiments are conducted using the mousetrap plug-ins for OpenSesame, the logging resolution can be specified explicitly in the experiment under "Logging resolution", which corresponds to the delay (in milliseconds) between recordings of the mouse position. By default, mouse positions are recorded every 10 ms (corresponding to a 100 Hz sampling rate). As the actual resolution achieved depends on the performance of the hardware, it makes sense to check the logging resolution using mt_check_resolution. Note that delays smaller than the specified delay typically result from mouse clicks in the experiment.

Value

A list with various descriptive statistics. For convenience, the relative frequencies are rounded to 4 decimal places.

Author(s)

Pascal J. Kieslich
Felix Henninger

Examples

mt_check_resolution(mt_example)

mt_cluster

Cluster trajectories.

Description

Performs trajectory clustering. It first computes distances between each pair of trajectories and then applies off-the-shelf clustering tools to explain the resulting dissimilarity matrix using a predefined number of clusters.
Usage

mt_cluster(
  data,
  use = "ln_trajectories",
  save_as = "clustering",
  dimensions = c("xpos", "ypos"),
  n_cluster = 5,
  method = "hclust",
  weights = rep(1, length(dimensions)),
  pointwise = TRUE,
  minkowski_p = 2,
  hclust_method = "ward.D",
  kmeans_nstart = 10,
  na_rm = FALSE,
  cluster_output = FALSE,
  verbose = FALSE
)

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use a character string specifying which trajectory data should be used.

save_as a character string specifying where the resulting data should be stored.

dimensions a character vector specifying which trajectory variables should be used. Can be of length 2 or 3, for two-dimensional or three-dimensional trajectories respectively.

n_cluster an integer specifying the number of clusters to estimate.

method character string specifying the clustering procedure. Either hclust (the default) or kmeans.

weights numeric vector specifying the relative importance of the variables specified in dimensions. Defaults to a vector of 1s implying equal importance. Technically, each variable is rescaled so that the standard deviation matches the corresponding value in weights. To use the original variables, set weights = NULL.

pointwise boolean specifying the way in which dissimilarity between the trajectories is measured. If TRUE (the default), mt_distmat measures the average dissimilarity and then sums the results. If FALSE, mt_distmat measures dissimilarity once (by treating the various points as independent dimensions). This is only relevant if method is "hclust". See mt_distmat for further details.

minkowski_p an integer specifying the distance metric for the cluster solution. minkowski_p = 1 computes the city-block distance, minkowski_p = 2 (the default) computes the Euclidian distance, minkowski_p = 3 the cubic distance, etc. Only relevant if method is "hclust". See mt_distmat for further details.

hclust_method character string specifying the linkage criterion used. Passed on to the method argument of hclust. Default is set to ward.D. Only relevant if method is "hclust".
kmeans_nstart  integer specifying the number of reruns of the kmeans procedure. Larger numbers minimize the risk of finding local minima. Passed on to the nstart argument of kmeans. Only relevant if method is "kmeans".

na_rm  logical specifying whether trajectory points containing NAs should be removed. Removal is done column-wise. That is, if any trajectory has a missing value at, e.g., the 10th recorded position, the 10th position is removed for all trajectories. This is necessary to compute distance between trajectories.

cluster_output  logical. If FALSE (the default), the mousetrap data object with the cluster assignments is returned (see Value). If TRUE, the output of the cluster method (kmeans or hclust) is returned directly.

verbose  logical indicating whether function should report its progress.

Details

mt_cluster uses off-the-shelf clustering tools, i.e., hclust and kmeans, for cluster estimation. Cluster estimation using hclust relies on distances computed by mt_distmat.

Mouse trajectories often occur in distinct, qualitative types (see Wulff et al., 2019; Wulff et al., 2021). Common trajectory types are linear trajectories, mildly and strongly curved trajectories, and single and multiple change-of-mind trials (see also mt_map). mt_cluster can tease these types apart.

mt_cluster uses hclust or kmeans to explain the distances between every pair of trajectories using a predefined number of clusters. If method is "hclust", mt_cluster computes the dissimiliarity matrix for all trajectory pairs using mt_distmat. If method is "kmeans", this is done internally by kmeans.

We recommend setting method to hclust using ward.D as the linkage criterion (via hclust_method). Relative to kmeans, the other implemented clustering method, and other linkage criteria, this setup handles the skewed distribution cluster sizes and trajectory outliers found in the majority of datasets best.

For clustering trajectories, it is often useful that the endpoints of all trajectories share the same direction, e.g., that all trajectories end in the top-left corner of the coordinate system (mt_remap_symmetric or mt_align can be used to achieve this). Furthermore, it is recommended to use length normalized trajectories (see mt_length_normalize; Wulff et al., 2019).

Value

A mousetrap data object (see mt_example) with an additional data.frame added to it (by default called clustering) that contains the cluster assignments. If a trajectory array was provided directly as data, only the clustering data.frame will be returned.

Author(s)

Dirk U. Wulff
Jonas M. B. Haslbeck
See Also

mt_distmat for more information about how the distance matrix is computed when the hclust method is used.

mt_cluster_k for estimating the optimal number of clusters.

Examples

# Length normalize trajectories
KH2017 <- mt_length_normalize(KH2017)

# Cluster trajectories
KH2017 <- mt_cluster(KH2017, use="ln_trajectories")

# Plot clustered trajectories
mt_plot(KH2017, use="ln_trajectories", use2="clustering", facet_col="cluster")

### mt_cluster_k

**Estimate optimal number of clusters.**

**Description**

Estimates the optimal number of clusters (k) using various methods.

**Usage**

mt_cluster_k(
  data,
  use = "ln_trajectories",
  dimensions = c("xpos", "ypos"),
  kseq = 2:15,
  compute = c("stability", "gap", "jump", "slope"),
  method = "hclust",
  weights = rep(1, length(dimensions)),
  pointwise = TRUE,
  minkowski_p = 2,
  hclust_method = "ward.D"
mt_cluster_k

```r
kmeans_nstart = 10,
n_bootstrap = 10,
model_based = FALSE,
n_gap = 10,
na_rm = FALSE,
verbose = FALSE
)
```

Arguments

data: a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use: a character string specifying which trajectory data should be used.

dimensions: a character vector specifying which trajectory variables should be used. Can be of length 2 or 3, for two-dimensional or three-dimensional trajectories respectively.

kseq: a numeric vector specifying set of candidates for k. Defaults to 2:15, implying that all values of k within that range are compared using the metrics specified in compute.

compute: character vector specifying the to be computed measures. Can be any subset of c("stability","gap","jump","slope").

method: character string specifying the type of clustering procedure for the stability-based method. Either hclust or kmeans.

weights: numeric vector specifying the relative importance of the variables specified in dimensions. Defaults to a vector of 1s implying equal importance. Technically, each variable is rescaled so that the standard deviation matches the corresponding value in weights. To use the original variables, set weights = NULL.

pointwise: boolean specifying the way in which dissimilarity between the trajectories is measured. If TRUE (the default), mt_distmat measures the average dissimilarity and then sums the results. If FALSE, mt_distmat measures dissimilarity once (by treating the various points as independent dimensions). This is only relevant if method is "hclust". See mt_distmat for further details.

minkowski_p: an integer specifying the distance metric for the cluster solution. minkowski_p = 1 computes the city-block distance, minkowski_p = 2 (the default) computes the Euclidian distance, minkowski_p = 3 the cubic distance, etc. Only relevant if method is "hclust". See mt_distmat for further details.

hclust_method: character string specifying the linkage criterion used. Passed on to the method argument of hclust. Default is set to ward.D. Only relevant if method is "hclust".

kmeans_nstart: integer specifying the number of reruns of the kmeans procedure. Larger numbers minimize the risk of finding local minima. Passed on to the nstart argument of kmeans. Only relevant if method is "kmeans".

n_bootstrap: an integer specifying the number of bootstrap comparisons used by stability. See cStability.
model_based  boolean specifying whether the model-based or the model-free should be used by stability, when method is kmeans. See cStability and Haslbeck & Wulff (2020).

n_gap  integer specifying the number of simulated datasets used by gap. See Tibshirani et al. (2001).

na_rm  logical specifying whether trajectory points containing NAs should be removed. Removal is done column-wise. That is, if any trajectory has a missing value at, e.g., the 10th recorded position, the 10th position is removed for all trajectories. This is necessary to compute distance between trajectories.

verbose  logical indicating whether function should report its progress.

Details

`mt_cluster_k` estimates the number of clusters (k) using four commonly used k-selection methods (specified via `compute`): cluster stability (`stability`), the gap statistic (`gap`), the jump statistic (`jump`), and the slope statistic (`slope`).

Cluster stability methods select k as the number of clusters for which the assignment of objects to clusters is most stable across bootstrap samples. This function implements the model-based and model-free methods described by Haslbeck & Wulff (2020). See references.

The remaining three methods select k as the value that optimizes the gap statistic (Tibshirani, Walther, & Hastie, 2001), the jump statistic (Sugar & James, 2013), and the slope statistic (Fujita, Takahashi, & Patriota, 2014), respectively.

For clustering trajectories, it is often useful that the endpoints of all trajectories share the same direction, e.g., that all trajectories end in the top-left corner of the coordinate system (mt_remap_symmetric or mt_align can be used to achieve this). Furthermore, it is recommended to use length normalized trajectories (see mt_length_normalize; Wulff et al., 2019).

Value

A list containing two lists that store the results of the different methods. `kopt` contains the estimated k for each of the methods specified in `compute`. `paths` contains the values for each k in `kseq` as computed by each of the methods specified in `compute`. The values in `kopt` are optima for each of the vectors in `paths`.

Author(s)

Dirk U. Wulff
Jonas M. B. Haslbeck

References


**See Also**

*mt_distmat* for more information about how the distance matrix is computed when the hclust method is used.

*mt_cluster* for performing trajectory clustering with a specified number of clusters.

**Examples**

```r
## Not run:
# Length normalize trajectories
KH2017 <- mt_length_normalize(KH2017)

# Find k
results <- mt_cluster_k(KH2017, use="ln_trajectories")

# Retrieve results
results$kopt
results$paths

## End(Not run)
```

---

**mt_count**

*Count number of observations.*

**Description**

Count number of observations per trial for a specified dimension (or several) in the trajectory array. This is mostly a helper function used by other functions in this package.

**Usage**

```r
mt_count(data, use = "trajectories", save_as = "measures", dimensions = "xpos")
```
Arguments

data: a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use: a character string specifying which trajectory data should be used.

save_as: a character string specifying where the resulting trajectory data should be stored.

dimensions: a character vector specifying the name of the dimension(s) that should be used for counting the number of observations. If several dimensions are specified, the number of complete observations are reported.

Value

A mousetrap data object (see mt_example).

If a data.frame with label specified in save_as (by default "measures") already exists, the number of observations (called nobs) are added as additional column. If not, an additional data.frame will be added.

If a trajectory array was provided directly as data, only a named character vector will be returned.

Author(s)

Pascal J. Kieslich

Examples

# Retrieve vector that counts number of observations
mt_count(mt_example$trajectories)

mt_derivatives

Calculate distance, velocity, and acceleration.

Description

Calculate distance traveled, velocity, and acceleration for each logged position. Distance is calculated as the Euclidean distance between successive coordinates, and velocity as distance covered per time interval. The acceleration denotes the difference in absolute velocity, again normalized per time.

Usage

mt_derivatives(
  data,
  use = "trajectories",
  save_as = use,
  dimensions = c("xpos", "ypos"),
  timestamps = "timestamps",
)
mt_derivatives

```r
prefix = "",
absolute = FALSE,
return_delta_time = FALSE,
verbose = FALSE
)
```

**Arguments**

- **data**: a mousetrap data object created using one of the mt_import functions (see `mt_example` for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).
- **use**: a character string specifying which trajectory data should be used.
- **save_as**: a character string specifying where the resulting trajectory data should be stored.
- **dimensions**: a character vector specifying across which dimension(s) distances, velocity, and acceleration are calculated. By default (`c("xpos","ypos")`), they are calculated across both x and y dimensions. Alternatively, only one dimension can be specified, e.g., "xpos" or "ypos".
- **timestamps**: a character string specifying the trajectory dimension containing the timestamps.
- **prefix**: an optional character string that is added as a prefix to the to be created new trajectory dimensions.
- **absolute**: logical indicating if absolute values for distances and velocities should be reported. Only relevant if a single dimension is specified in `dimensions` (see Details).
- **return_delta_time**: logical indicating if the timestamp differences should be returned as well (as "delta_time").
- **verbose**: logical indicating whether function should report its progress.

**Details**

Distances, velocities and acceleration are computed as follows:

The first entry in each respective vector is always zero. Each subsequent entry thus represents the Euclidean distance traveled since the previous recorded set of coordinates and the velocity with which the movement between both samples took place. Thus, both distance and velocity represent the intervening period between the previous sample and the one with which the numeric value is saved.

The acceleration, by contrast, denotes the change in absolute velocity between two adjacent periods. Because of this, it is shifted forward to best match the actual time point at which the acceleration was measured. Because there will always be one less value computed for acceleration than for velocity, the final value in the acceleration vector has been padded with an NA.

If the distance is calculated across both horizontal and vertical (x and y) dimensions, distance and velocity is always positive (or 0). If only one dimension is used, by default (absolute=FALSE), increases in x (or y) values result in positive distances and velocity values, decreases in negative distances and velocity values. If absolute=TRUE, absolute values for distance and velocity are reported.
Value

A mousetrap data object (see mt_example) with Euclidian distance, velocity, and acceleration added as additional variables to the trajectory array (called dist, vel, and acc, if no prefix was specified). If the trajectory array was provided directly as data, only the trajectory array will be returned.

Author(s)

Pascal J. Kieslich
Felix Henninger

See Also

mt_average for averaging trajectories across constant time intervals.
mt_measures for calculating per-trial mouse-tracking measures.

Examples

```r
# Calculate derivatives looking at movement
# across both dimensions
mt_example <- mt_derivatives(mt_example)

# Calculate derivatives only looking at movement along x dimension
# reporting absolute values for distance and velocity
mt_example <- mt_derivatives(mt_example,
  dimensions="xpos", absolute=TRUE)
```

Description

Calculate the idealized trajectory and the perpendicular deviations of the actual trajectory from it for each logged position.

Usage

```r
mt_deviations(
  data,
  use = "trajectories",
  save_as = use,
  dimensions = c("xpos", "ypos"),
  start_ideal = NULL,
  end_ideal = NULL,
  prefix = "",
  verbose = FALSE
)
```
**Arguments**

- **data**: a mousetrap data object created using one of the `mt_import` functions (see `mt_example` for details). Alternatively, a trajectory array can be provided directly (in this case `use` will be ignored).
- **use**: a character string specifying which trajectory data should be used.
- **save_as**: a character string specifying where the resulting trajectory data should be stored.
- **dimensions**: a character vector specifying the two dimensions in the trajectory array that contain the mouse positions. By default (c("xpos","ypos")), the x- and y-positions are used.
- **start_ideal**: an optional vector specifying the start position (see Example). If specified, this position will be used as the starting point of the idealized trajectory (instead of the actual starting point).
- **end_ideal**: an optional vector specifying the end position (see Example). If specified, this position will be used as the end point of the idealized trajectory (instead of the actual end point).
- **prefix**: an optional character string that is added as a prefix to the to be created new trajectory dimensions.
- **verbose**: logical indicating whether function should report its progress.

**Details**

The idealized trajectory is defined as the straight line connecting the start and end point of the actual trajectory (e.g., Freeman & Ambady, 2010). The deviation for each position is calculated as the perpendicular deviation of the actual trajectory from the idealized trajectory.

If a deviation occurs above the direct path, this is denoted by a positive value. If it occurs below the direct path, this is denoted by a negative value. This assumes that the complete movement in the trial was from bottom to top (i.e., the end point has a higher y-position than the start points). In case the movement was from top to bottom, `mt_deviations` automatically flips the signs. Note that the second dimension specified in `dimensions` is used for determining all this.

**Value**

A mousetrap data object (see `mt_example`) where the positions of the idealized trajectory (by default called `xpos_ideal` and `ypos_ideal`) and the perpendicular deviations of the actual trajectory from the idealized trajectory (by default called `dev_ideal`) have been added as additional variables to the trajectory array. If the trajectory array was provided directly as `data`, only the trajectory array will be returned.

**Author(s)**

Pascal J. Kieslich  
Felix Henninger

**References**

See Also

mt_measures for calculating per-trial mouse-tracking measures.

Examples

# Calculate deviations from idealized trajectory
# (straight line connecting the start and end point of each trial)
mt_example <- mt_deviations(mt_example)

# Calculate deviations from idealized trajectory with
# constant start and end points across trials
mt_example <- mt_deviations(mt_example,
    start_ideal=c(0,0), end_ideal=c(-665,974))

Description

[Experimental]

mt_diffmap creates a difference-heatmap of the trajectory data using gaussian smoothing. Note that this function has beta status.

Usage

mt_diffmap(
    x,
    y = NULL,
    condition = NULL,
    use = "trajectories",
    dimensions = c("xpos", "ypos"),
    use2 = "data",
    filename = NULL,
    bounds = NULL,
    xres = 500,
    upscale = 4,
    smooth_radius = 10,
    colors = c("#00863F", "#000000", "#FF1900"),
    n_shades = 1000,
    plot = TRUE,
    ...
)

mt_diffmap

Creates a difference-heatmap of two trajectory heatmap images.
mt_diffmap

Arguments

\textbf{x} \hspace{1cm} \text{an object of class \texttt{mousetrap}}, a trajectory object of class \texttt{array}, or an object of class \texttt{mt_heatmap_raw} (as created by \texttt{mt_heatmap_raw}).

\textbf{y} \hspace{1cm} \text{an object of class \texttt{mousetrap}}, a trajectory object of class \texttt{array}, or an object of class \texttt{mt_heatmap_raw} (as created by \texttt{mt_heatmap_raw}). The class of \texttt{y} must match the class of \texttt{x}, unless \texttt{y} is \texttt{NULL}.

\textbf{condition} \hspace{1cm} \text{either a character value specifying which variable codes the two conditions (in \texttt{x[[use2]]}) that should be compared - or a vector matching the number of trajectories in \texttt{x[[use]]} that has exactly two levels. \texttt{mt_diffmap} will create a difference-heatmap comparing all trajectories between the two conditions. If condition is specified, \texttt{y} will be ignored (unless \texttt{x} and \texttt{y} are of class \texttt{heatmap_raw}).

\textbf{use} \hspace{1cm} \text{a character string specifying which trajectory data should be used.}

\textbf{dimensions} \hspace{1cm} \text{a character vector specifying the trajectory variables used to create the heatmap. The first two entries are used as x and y-coordinates, the third, if provided, will be added as color information.}

\textbf{use2} \hspace{1cm} \text{an optional character string specifying where the data that contain the condition variable can be found. Defaults to "data" as \texttt{x["data"]} usually contains all non mouse-tracking trial data.}

\textbf{filename} \hspace{1cm} \text{a character string giving the name of the file. If \texttt{NULL} (the default), the R standard device is used for plotting. Otherwise, the plotting device is inferred from the file extension. Only supports devices \texttt{tiff}, \texttt{png}, \texttt{pdf}.}

\textbf{bounds} \hspace{1cm} \text{numeric vector specifying the corners (xmin, ymin, xmax, ymax) of the plot region. By default (\texttt{bounds = NULL}), bounds are determined based on the data input.}

\textbf{xres} \hspace{1cm} \text{an integer specifying the number of pixels along the x-dimension. An \texttt{xres} of 1000 implies an 1000*N px, where N is determined so that the trajectories aspect ratio is preserved (provided the \texttt{bounds} are unchanged).}

\textbf{upscale} \hspace{1cm} \text{a numeric value by which the output resolution of the image is increased or decreased. Only applies if device is one of \texttt{tiff}, \texttt{png}, or \texttt{pdf}.}

\textbf{smooth_radius} \hspace{1cm} \text{a numeric value specifying the standard deviation of the gaussian smoothing. If zero, smoothing is omitted.}

\textbf{colors} \hspace{1cm} \text{a character vector specifying the colors used to color cases of \texttt{image1 > image2}, \texttt{image1 \sim image2}, \texttt{image1 < image2}, respectively. Note that the colors are used in that specific order. Defaults to \texttt{c("#00863F", "#FFFFFF", "#FF1900")} which specifies a green-black-red color gradient.}

\textbf{n_shades} \hspace{1cm} \text{integer specifying the number of shades for the color gradient between the first and second, and the second and third color in \texttt{colors}.}

\textbf{plot} \hspace{1cm} \text{logical specifying whether resulting image should be plotted (\texttt{plot = TRUE}, the default). If (\texttt{plot = FALSE}), an object of class \texttt{mt_object_raw} is returned.}

\textbf{...} \hspace{1cm} \text{arguments passed to \texttt{mt_heatmap_raw}.}

\textbf{verbose} \hspace{1cm} \text{logical indicating whether function should report its progress.}
Details

mt_diffmap takes two objects that either contain trajectory heatmaps or from which trajectory heatmaps can be computed. Difference-heatmaps are constructed analogously to mt_heatmap_raw.

Author(s)

Dirk U. Wulff
Pascal J. Kieslich

References


See Also

mt_heatmap and mt_heatmap_ggplot for plotting trajectory heatmaps.

Examples

mt_diffmap(KH2017, condition="Condition",
           xres=400, smooth_radius=6, n_shades=5)

mt_distmat

Compute distance matrix.

Description

Computes the point- or vector-wise dissimilarity between each pair of trajectories.

Usage

mt_distmat(
  data,
  use = "ln_trajectories",
  save_as = "distmat",
  dimensions = c("xpos", "ypos"),
  weights = rep(1, length(dimensions)),
  pointwise = TRUE,
  minkowski_p = 2,
  na_rm = FALSE
)
Arguments

data
a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use
a character string specifying which trajectory data should be used.

save_as
a character string specifying where the resulting data should be stored.

dimensions
a character vector specifying which trajectory variables should be used. Can be of length 2 or 3 for two-dimensional or three-dimensional trajectories respectively.

weights
numeric vector specifying the relative importance of the variables specified in dimensions. Defaults to a vector of 1s implying equal importance. Technically, each variable is rescaled so that the standard deviation matches the corresponding value in weights. To use the original variables, set weights = NULL.

pointwise
boolean specifying the way dissimilarity between the trajectories is measured (see Details). If TRUE (the default), mt_distmat measures the average dissimilarity and then sums the results. If FALSE, mt_distmat measures dissimilarity once (by treating the various points as independent dimensions).

minkowski_p
an integer specifying the distance metric. minkowski_p = 1 computes the city-block distance, minkowski_p = 2 (the default) computes the Euclidian distance, minkowski_p = 3 the cubic distance, etc.

na_rm
logical specifying whether trajectory points containing NAs should be removed. Removal is done column-wise. That is, if any trajectory has a missing value at, e.g., the 10th recorded position, the 10th position is removed for all trajectories. This is necessary to compute distance between trajectories.

Details

mt_distmat computes point- or vector-wise dissimilarities between pairs of trajectories. Point-wise dissimilarity refers to computing the distance metric defined by minkowski_p for every point of the trajectory and then summing the results. That is, if minkowski_p = 2 the point-wise dissimilarity between two trajectories, each defined by a set of x and y coordinates, is calculated as sum(sqrt((x_i-x_j)^2 + (y_i-y_j)^2)). Vector-wise dissimilarity, on the other hand refers to computing the distance metric once for the entire trajectory. That is, vector-wise dissimilarity is computed as sqrt(sum((x_i-x_j)^2 + (y_i-y_j)^2)).

Value

A mousetrap data object (see mt_example) with an additional object added (by default called distmat) containing the distance matrix. If a trajectory array was provided directly as data, only the distance matrix will be returned.

Author(s)

Dirk U. Wulff
Jonas M. B. Haslbeck
Examples

```r
# Length normalize trajectories
mt_example <- mt_length_normalize(mt_example)

# Compute distance matrix
mt_example <- mt_distmat(mt_example, use="ln_trajectories")
```

---

**mt_example**

_A mousetrap data object._

---

### Description

A data object of class "mousetrap" with example data created by importing `mt_example_raw` and applying basic post-processing.

### Usage

```r
mt_example
```

### Format

A mousetrap data object is a _list_ containing at least the following objects:

- _data_: a _data.frame_ containing the trial data (from which the mouse-tracking data columns have been removed). More information about the content of the trial data in `mt_example` can be found in `mt_example_raw`. The _rownames_ of _data_ correspond to the trial identifier. For convenience, the trial identifier is also stored in an additional column called "mt_id".

- _trajectories_: an _array_ containing the raw mouse-tracking trajectories. The first dimension represents the different trials and the dimension names (which can be accessed using _rownames_) correspond to the trial identifier (the same identifier that is used as the _rownames_ in _data_). The second dimension corresponds to the samples taken over time which are included in chronological order. The third dimension corresponds to the different mouse-tracking variables (timestamps, x-positions, y-positions) which are usually called _timestamps_, _xpos_, and _ypos_.

Some functions in this package (e.g., `mt_time_normalize` and `mt_average`) add additional trajectory arrays (e.g., _tn_trajectories_ and _av_trajectories_) to the mousetrap data object. Other functions modify the existing arrays (e.g., `mt_derivatives` adds distance, velocity, and acceleration to an existing dataset). Finally `mt_measures` adds an additional data.frame with mouse-tracking measures to it.

### Details

The raw data set was imported using `mt_import_mousetrap`. Trajectories were then remapped using `mt_remap_symmetric` so that all trajectories end in the top-left corner and their starting point was aligned using `mt_align_start` to a common value (0,0).
mt_example_raw

---

Description

An exemplary mouse-tracking dataset collected OpenSesame using the mousetrap plugin (Kieslich & Henninger, 2017). A preprocessed (as opposed to raw) version of the same data can be found in mt_example.

Usage

mt_example_raw

Format

A data.frame with 38 rows and 19 variables. The data.frame is based on the combined raw data that were created using read_opensesame from the readbulk library. For ease of use, unnecessary columns were excluded.

The variables included relate to the item that was presented (Exemplar), the answer categories (Category1 and Category2), the subject identifier (subject_nr) the subjects’ response (response_get_response), as well as the mouse-tracking variables (timestamps_get_response, xpos_get_response and ypos_get_response). Besides, a number of additional variables are included, e.g., some variables relating to the general settings of the experiment (e.g., the width and height of the screen in pixels).

Each mouse-tracking variable contains a list of values (separated by ', ')

- one entry for each recorded position of the mouse. The position coordinates are given in pixels, such that values of zero for both xpos_get_response and ypos_get_response indicate that the cursor is located in the center of the screen. Both variables increase in value as the mouse moves toward the bottom right. Timestamps are given in milliseconds.

Details

The data stem from a study based on experiment 1 by Dale et al. (2007). In this experiment, participants have to assign exemplars (e.g., "shark") to one of two categories (e.g., "fish" or "mammal") by clicking on the button corresponding to the correct category. All exemplars and categories were translated to and presented in German.

Across the 19 trials of the experiment, participants categorized 13 exemplars that were typical of their category and 6 atypical exemplars for which this was not the case. For the atypical exemplars (e.g., "whale"), the competing category ("fish") was selected to compete with the correct category ("mammal"). The hypothesis under investigation is whether participants’ mouse trajectories deviate more towards the competing category for the atypical exemplars, indicating increased conflict between the response options.

Please note that mt_example_raw should only be used for exploring the features of the mousetrap package and not for any substantive analysis.
References


---

mt_exclude_finish  
*Exclude phase without mouse movement at end of trial.*

Description

Exclude a potential phase at the end of a trial where the mouse was not moved. The corresponding samples (x- and y-positions and timestamps) in the trajectory data will be removed.

Usage

``` r
mt_exclude_finish(
  data,
  use = "trajectories",
  save_as = use,
  dimensions = c("xpos", "ypos"),
  timestamps = "timestamps",
  verbose = FALSE
)
```

Arguments

- **data**: a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).
- **use**: a character string specifying which trajectory data should be used.
- **save_as**: a character string specifying where the resulting trajectory data should be stored.
- **dimensions**: a character vector specifying the dimensions in the trajectory array that contain the mouse positions.
- **timestamps**: a character string specifying the trajectory dimension containing the timestamps.
- **verbose**: logical indicating whether function should report its progress.

Details

`mt_exclude_finish` removes all samples (except the first) at the end of the trial during which the mouse was not moved compared to its final position. It returns only x- and y-positions as well as timestamps.

Please note that this operation may result in changes in several mouse-tracking measures, for example, the response time (RT).
Value
A mousetrap data object (see mt_example) from which a potential phase without mouse movement at the end of the trial was removed. If the trajectory array was provided directly as data, only the trajectory array will be returned.

Author(s)
Pascal J. Kieslich
Dirk U. Wulff

See Also
mt_exclude_initiation for removing a potential initial phase without mouse movement.

Examples
```r
mt_example <- mt_exclude_finish(mt_example,
  save_as="mod_trajectories")
```

Description
Exclude the initial phase in a trial where the mouse was not moved. The corresponding samples (x- and y-positions and timestamps) in the trajectory data will be removed.

Usage
```r
mt_exclude_initiation(
  data, 
  use = "trajectories", 
  save_as = use, 
  dimensions = c("xpos", "ypos"), 
  timestamps = "timestamps", 
  reset_timestamps = TRUE, 
  verbose = FALSE
)
```

Arguments
- **data**: a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).
- **use**: a character string specifying which trajectory data should be used.
- **save_as**: a character string specifying where the resulting trajectory data should be stored.
mt_exclude_initiation

dimensions a character vector specifying the dimensions in the trajectory array that contain the mouse positions.
timestamps a character string specifying the trajectory dimension containing the timestamps.
reset_timestamps logical indicating whether the timestamps should be reset after removing the initial phase without movement (see Details).
verbose logical indicating whether function should report its progress.

Details

mt_exclude_initiation removes all samples (x- and y-positions as well as timestamps) at the beginning of the trial during which the mouse was not moved from its initial position. The last unchanged sample is retained in the data.

If reset_timestamps == TRUE (the default), it subtracts the last timestamp before a movement occurs from all timestamps, so that the series of timestamps once more begin with zero. If the argument is set to FALSE, the values of the timestamps are unchanged.

Please note that resetting the timestamps will result in changes in several mouse-tracking measures, notably those which report timestamps (e.g., MAD_time). Typically, however, these changes are desired when using this function.

Value

A mousetrap data object (see mt_example) from which the initial phase without mouse movement was removed. If the trajectory array was provided directly as data, only the trajectory array will be returned.

Author(s)

Pascal J. Kieslich
Felix Henninger

See Also

mt_measures for calculating the initiation time.

mt_exclude_finish for removing a potential phase without mouse movement at the end of the trial.

Examples

mt_example <- mt_exclude_initiation(mt_example,
save_as="mod_trajectories")
Export mouse-tracking data.

Description

`mt_export_long` and `mt_export_wide` can be used for exporting mouse-tracking data from a mousetrap data object in long or wide format. If desired, additional data (stored in `data[[use2]]`) can be merged with the trajectory data before export. `mt_export_long` and `mt_export_wide` are wrapper functions for `mt_reshape`.

Usage

```r
mt_export_long(
  data,
  use = "trajectories",
  use_variables = NULL,
  use2 = "data",
  use2_variables = NULL,
  ...
)
```

```r
mt_export_wide(
  data,
  use = "trajectories",
  use_variables = NULL,
  use2 = "data",
  use2_variables = NULL,
  ...
)
```

Arguments

- **data**: a mousetrap data object created using one of the `mt_import` functions (see `mt_example` for details). Alternatively, a trajectory array can be provided directly (in this case `use` will be ignored).
- **use**: a character string specifying which data should be exported. The corresponding data are selected from data using `data[[use]]`. Usually, this value corresponds to either "trajectories" or "tn_trajectories", depending on whether the raw or the time-normalized trajectories should be exported.
- **use_variables**: a character vector specifying which mouse-tracking variables should be exported. This corresponds to the labels of the trajectory array dimensions. If unspecified, all variables will be exported.
- **use2**: an optional character string specifying where the other trial data can be found. Defaults to "data" as `data[["data"]]]` usually contains all non mouse-tracking trial data. Alternatively, a data.frame can be provided directly.
use2_variables  an optional character string (or vector) specifying the variables (in data[[use2]]) that should be merged with the data.

... additional arguments passed on to mt reshape (such as subset).

Value

A data.frame containing the exported data.

Functions

• mt_export_long: Export mouse-tracking data in long format
• mt_export_wide: Export mouse-tracking data in wide format

Author(s)

Pascal J. Kieslich
Felix Henninger

See Also

mt_import_long for importing mouse-tracking data saved in a long format.
mt_import_wide for importing mouse-tracking data saved in a wide format.

Examples

# Export data in long format
# (and include information about condition and subject_nr)
mt_data_long <- mt_export_long(mt_example,
    use2_variables=c("subject_nr","Condition"))

# Export data in wide format
# (and include information about condition and subject_nr)
mt_data_wide <- mt_export_wide(mt_example,
    use2_variables=c("subject_nr","Condition"))

mt_heatmap  Plot trajectory heatmap.

Description

[Experimental]

mt_heatmap plots high resolution raw trajectory maps. Note that this function has beta status.
Usage

mt_heatmap(
  x,
  use = "trajectories",
  dimensions = c("xpos", "ypos"),
  filename = NULL,
  ..., 
  upscale = 1,
  plot_dims = FALSE,
  verbose = TRUE
)

Arguments

x usually an object of class mousetrap. Alternatively, a trajectory array or an object of class mt_heatmap_raw.
use a character string specifying which trajectory data should be used.
dimensions a character vector specifying the trajectory variables used to create the heatmap. The first two entries are used as x and y-coordinates, the third, if provided, will be added as color information.
filename a character string giving the name of the file. If NULL (the default), the R standard device is used for plotting. Otherwise, the plotting device is inferred from the file extension. Only supports devices tiff, png, pdf.
...
arguments passed to mt_heatmap_raw.
upscale a numeric value by which the output resolution of the image is increased or decreased. Only applies if device is one of tiff, png, or pdf.
plot_dims adds the coordinates of the four image corners to the plot. Helps setting bounds.
verbose logical indicating whether function should report its progress.

Details

mt_heatmap wraps mt_heatmap_raw and provides direct plotting output in tiff, png, pdf, or R’s default window output. For further details on how the trajectory heatmaps are constructed, see mt_heatmap_raw.

Author(s)

Dirk U. Wulff

References


See Also

mt_heatmap_ggplot for plotting a trajectory heatmap using ggplot2.
mt_diffmap for plotting trajectory difference-heatmaps.

Examples

mt_heatmap(KH2017, xres=500, n_shades=5, mean_image=0.2)

mt_heatmap_ggplot

Plot trajectory heatmap using ggplot.

Description

[Experimental]

mt_heatmap_ggplot plots high resolution raw trajectory maps. Note that this function has beta status.

Usage

mt_heatmap_ggplot(
  data,
  use = "trajectories",
  dimensions = c("xpos", "ypos"),
  use2 = "data",
  facet_row = NULL,
  facet_col = NULL,
  ...)

Arguments

data : a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use : a character string specifying which trajectory data should be used.

dimensions : a character vector specifying the trajectory variables used to create the heatmap. The first two entries are used as x and y-coordinates, the third, if provided, will be added as color information.

use2 : an optional character string specifying where the data that contain the variables used for faceting can be found (in case these arguments are specified). Defaults to "data" as data[["data"]]) usually contains all non mouse-tracking trial data.
mt_heatmap_ggplot

facet_row  an optional character string specifying a variable in data[[use2]] that should be used for (row-wise) faceting.

facet_col  an optional character string specifying a variable in data[[use2]] that should be used for (column-wise) faceting.

... arguments passed to mt_heatmap_raw.

Details

mt_heatmap_ggplot wraps mt_heatmap_raw and returns a ggplot object containing the plot. In contrast to mt_heatmap_plot plots created by mt_heatmap_ggplot can be extended using ggplot’s + operator. For further details on how the trajectory heatmaps are constructed, see mt_heatmap_raw.

Author(s)

Pascal J. Kieslich
Felix Henninger
Dirk U. Wulff

References


See Also

mt_heatmap for plotting a trajectory heatmap using base plots.
mt_diffmap for plotting trajectory difference-heatmaps.

Examples

mt_heatmap_ggplot(KH2017, xres=500, n_shades=5, mean_image=0.2)
mt_heatmap_raw  

Creates high-resolution heatmap of trajectory data.

Description

[Experimental]

mt_heatmap_raw creates a high-resolution heatmap image of the trajectory data using gaussian smoothing. Note that this function has beta status.

Usage

mt_heatmap_raw(  
data,  
use = "trajectories",  
dimensions = c("xpos", "ypos"),  
variable = NULL,  
bounds = NULL,  
xres = 1000,  
upsample = 1,  
norm = FALSE,  
colors = c("black", "blue", "white"),  
n_shades = c(1000, 1000),  
smooth_radius = 1.5,  
low_pass = 200,  
auto_enhance = TRUE,  
mean_image = 0.15,  
mean_color = 0.25,  
aggregate_lwd = 0,  
aggregate_col = "black",  
n_trajectories = NULL,  
seed = NULL,  
verbose = TRUE  
)

Arguments

data  
a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use  
a character string specifying which trajectory data should be used.

dimensions  
a character vector specifying the trajectory variables used to create the heatmap. The first two entries are used as x and y-coordinates, the third, if provided, will be added as color information.

variable  
boolean or numeric vector matching the number of trajectories that if provided will be used as color information. variable is only considered when length(dimensions) < 3.
### Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>bounds</td>
<td>numeric vector specifying the corners (xmin, ymin, xmax, ymax) of the plot region. By default (bounds = NULL), bounds are determined based on the data input.</td>
</tr>
<tr>
<td>xres</td>
<td>an integer specifying the number of pixels along the x-dimension. An xres of 1000 implies an 1000*N px, where N is determined so that the trajectories aspect ratio is preserved (provided the bounds are unchanged).</td>
</tr>
<tr>
<td>upsample</td>
<td>a numeric value by which the number of points used to represent individual trajectories are increased or decreased. Values of smaller than one will improve speed but also introduce a certain level of granularity.</td>
</tr>
<tr>
<td>norm</td>
<td>a logical specifying whether the data should be warped into standard space. If norm = TRUE, this overrules bounds.</td>
</tr>
<tr>
<td>colors</td>
<td>a character vector specifying two or three colors used to color the background, the foreground (trajectories), and the values of a third dimension (if specified).</td>
</tr>
<tr>
<td>n_shades</td>
<td>an integer specifying the number of shades for the color gradient between the first and second, and the second and third color in colors.</td>
</tr>
<tr>
<td>smooth_radius</td>
<td>a numeric value specifying the standard deviation of the gaussian smoothing. If zero, smoothing is omitted.</td>
</tr>
<tr>
<td>low_pass</td>
<td>an integer specifying the allowed number of counts per pixel. This arguments limits the maximum pixel color intensity.</td>
</tr>
<tr>
<td>auto_enhance</td>
<td>boolean. If TRUE (the default), the image is adjusted so that the mean color intensity matches mean_image and mean_color.</td>
</tr>
<tr>
<td>mean_image</td>
<td>a numeric value between 0 and 1 specifying the average foreground color intensity across the entire image. Defaults to 0.1.</td>
</tr>
<tr>
<td>mean_color</td>
<td>a numeric value between 0 and 1 specifying the average third dimension’s color intensity across the entire image. Defaults to 0.1. Only relevant if a third dimension is specified in colors.</td>
</tr>
<tr>
<td>aggregate_lwd</td>
<td>an integer specifying the width of the aggregate trajectory. If aggregate_lwd is 0 (the default), the aggregate trajectory is omitted.</td>
</tr>
<tr>
<td>aggregate_col</td>
<td>a character value specifying the color of the aggregate trajectory.</td>
</tr>
<tr>
<td>n_trajectories</td>
<td>an optional integer specifying the number of trajectories used to create the image. By default, all trajectories are used. If n_trajectories is specified and smaller than the number of trajectories in the trajectory array, then n_trajectories are randomly sampled.</td>
</tr>
<tr>
<td>seed</td>
<td>an optional integer specifying the seed used for the trajectory sampling.</td>
</tr>
<tr>
<td>verbose</td>
<td>logical indicating whether function should report its progress.</td>
</tr>
</tbody>
</table>

### Details

To create the image, `mt_heatmap_raw` takes the following steps. First, the function maps the trajectory points to a pixel space with x ranging from 1 to xres and y ranging from 1 to xres divided by the ratio of x and y’s value range. Second, the function counts and normalizes the number of trajectory points occupying each of the x,y-pixels to yield image intensities between 0 and 1. Third, the function smooths the image using an approximative guassian approach governed by smooth_radius,
which controls the dispersion of the gaussian smoothing. Fourth, the function automatically enhances the image (unless auto_enhance = FALSE) using a non-linear transformation in order to yield a desired mean_image intensity. Fifth, the function translates the image intensity into color using the colors specified in colors. Finally, the function returns the image data in a long format containing the x, y, and color information.

`mt_heatmap_raw` also offers the possibility to overlay the heatmap with an additional variable, such as for instance velocity, so that both the density of mouse trajectories and the information of the additional variable are visible. In order to do this, specify a third variable label in dimensions and control its appearance using the color and mean_color arguments.

Value

An object of class `mt_object_raw` containing in a matrix format the image’s pixel information, the aggregate trajectory, and the colors.

Author(s)

Dirk U. Wulff

References


See Also

`mt_heatmap` and `mt_heatmap_ggplot` for plotting trajectory heatmaps.

`mt_diffmap` for plotting trajectory difference-heatmaps.

**mt_import_long**

*Import mouse-tracking data saved in long format.*

Description

`mt_import_long` receives a data.frame in which mouse-tracking data are stored in long format, i.e., where one row contains the logging data (timestamp, x- and y-position etc.) at one specific point in the trial. This is, for example, the case when exporting the trajectory data from the mousetrap package using `mt_export_long`. From this data.frame, `mt_import_long` creates a mousetrap data object containing the trajectories and additional data for further processing within the mousetrap package. Specifically, it returns a list that includes the trajectory data as an array (called trajectories), and all other data as a data.frame (called data). This data structure can then be passed on to other functions within this package (see mousetrap for an overview).
### Usage

```r
mt_import_long(
  raw_data,
  xpos_label = "xpos",
  ypos_label = "ypos",
  zpos_label = NULL,
  timestamps_label = "timestamps",
  add_labels = NULL,
  mt_id_label = "mt_id",
  mt_seq_label = "mt_seq",
  reset_timestamps = TRUE,
  verbose = TRUE
)
```

### Arguments

- `raw_data`: a data.frame in long format, containing the raw data.
- `xpos_label`: a character string specifying the column containing the x-positions.
- `ypos_label`: a character string specifying the column containing the y-positions.
- `zpos_label`: an optional character string specifying the column containing the z-positions.
- `timestamps_label`: a character string specifying the column containing the timestamps. If no timestamps are found in the data, a timestamps variable with increasing integers will be created (assuming equidistant time steps).
- `add_labels`: a character vector specifying columns containing additional mouse-tracking variables.
- `mt_id_label`: a character string (or vector) specifying the name of the column that provides a unique ID for every trial (the trial identifier). If more than one variable name is provided, a new ID variable will be created by combining the values of each variable. The trial identifier will be set as the rownames of the resulting trajectories and trial data, and additionally be stored in the column "mt_id" in the trial data.
- `mt_seq_label`: a character string specifying the column that indicates the order of the logged coordinates within a trial. If no column of the specified name is found in the data.frame, the coordinates will be imported in the order in which they were stored in `raw_data`.
- `reset_timestamps`: logical indicating if the first timestamp should be subtracted from all timestamps within a trial. Default is `TRUE` as it is recommended for all following analyses in mousetrap.
- `verbose`: logical indicating whether function should report its progress.

### Details

The default arguments are set so that no adjustments have to be made when importing a data.frame that was created using `mt_export_long`. 
The coordinates are ordered according to the values in the column provided in the `mt_seq_label` parameter (default `mt_seq`). If the corresponding column does not exist, the coordinates will be imported in the order in which they were stored in the raw data.

If no timestamps are found in the data, `mt_import_long` automatically creates a timestamps variable with increasing integers (starting with 0) assuming equally spaced sampling intervals.

**Value**

A mousetrap data object (see `mt_example`).

**Author(s)**

Pascal J. Kieslich
Felix Henninger

**See Also**

`mt_import_mousetrap` and `mt_import_wide` for importing mouse-tracking data in other formats.

**Examples**

```r
# Create data in long format for test purposes
mt_data_long <- mt_export_long(mt_example,
   use2_variables=c("subject_nr","Condition"))

# Import the data using mt_import_long
mt_data <- mt_import_long(mt_data_long)
```

```r
## Not run:
# Import a hypothetical dataset that contains the
# custom mouse-tracking variables angle and velocity
mt_data <- mt_import_long(exp_data,
   add_labels= c("angle", "velocity"))

## End(Not run)
```

---

**mt_import_mousetrap**

Import mouse-tracking data recorded using the mousetrap plug-ins in OpenSesame.

**Description**

`mt_import_mousetrap` accepts a data.frame of (merged) raw data from a mouse-tracking experiment implemented in OpenSesame using the mousetrap plugin (Kieslich & Henninger, 2017). From this data.frame, `mt_import_mousetrap` creates a mousetrap data object containing the trajectories and additional data for further processing within the mousetrap package. Specifically, it returns
a list that includes the trajectory data as an array (called trajectories), and all other data as a data.frame (called data). This data structure can then be passed on to other functions within this package (see mousetrap for an overview).

Usage

mt_import_mousetrap(
  raw_data,
  xpos_label = "xpos",
  ypos_label = "ypos",
  timestamps_label = "timestamps",
  mt_id_label = NULL,
  split = ",",
  duplicates = "remove_first",
  unordered = "warn",
  reset_timestamps = TRUE,
  digits = NULL,
  verbose = FALSE
)

Arguments

raw_data a data.frame containing the raw data.
xpos_label a character string specifying the name of the column(s) in which the x-positions are stored (see Details).
ypos_label a character string specifying the name of the column(s) in which the y-positions are stored (see Details).
timestamps_label a character string specifying the name of the column(s) in which the timestamps are stored (see Details).
mt_id_label an optional character string (or vector) specifying the name of the column that provides a unique ID for every trial (the trial identifier). If unspecified (the default), an ID variable will be generated. If more than one variable name is provided, a new ID variable will be created by combining the values of each variable. The trial identifier will be set as the rownames of the resulting trajectories and trial data, and additionally be stored in the column "mt_id" in the trial data.
split a character string indicating how the different timestamps and coordinates within a trial are separated.
duplicates a character string indicating how duplicate timestamps within a trial are handled (see Details).
unordered a character string indicating how unordered (i.e., non-monotonically increasing) timestamps within a trial are handled (see Details).
reset_timestamps logical indicating if the first timestamp should be subtracted from all timestamps within a trial. Default is TRUE as it is recommended for all following analyses in mousetrap.
\texttt{mt_import_mousetrap}

digits\hspace{1cm} an optional integer. If specified, timestamps will be rounded. Potentially useful if timestamps are recorded with submillisecond precision.

verbose\hspace{1cm} logical indicating whether function should report its progress.

\textbf{Details}

When working with mouse-tracking data that were recorded using the mousetrap plug-ins for OpenSesame, usually only the \texttt{raw\_data} need to be provided. All other arguments have sensible defaults.

If the relevant timestamps, x-positions, and y-positions are each stored in one variable, a character string specifying (parts of) the respective column name needs to be provided. In this case, the column names are extracted using \texttt{grep} to find the column that starts with the respective character string (in OpenSesame these will typically contain the name of the item that was used to record them, such as \texttt{xpos\_get\_response}). This means that the exact column names do not have to be provided - as long as only one column starts with the respective character string (otherwise, the exact column names have to be provided).

If several variables contain the timestamps, x-positions, and y-positions within a trial (e.g., \texttt{xpos\_part1} and \texttt{xpos\_part2}), a vector of the exact column names has to be provided (e.g., \texttt{xpos\_label=c(\"xpos\_part1\", \"xpos\_part2\")}). \texttt{mt_import_mousetrap} will then merge all raw data in the order with which the variable labels have been specified. If one variable contains NAs or an empty string in a trial, these cases will be ignored (this covers the special case that, e.g., \texttt{xpos\_part2} is only relevant for some trials and contains NAs in the other trials).

duplicates allows for different options to handle duplicate timestamps within a trial:

- remove\_first: First timestamp and corresponding x-/y-positions are removed (the default).
- remove\_last: Last timestamp and corresponding x-/y-positions are removed.
- ignore: Duplicates are kept.

unordered allows for different options to handle unordered, that is, non-monotonically increasing timestamps within a trial:

- warn: A warning is issued if unordered timestamps are encountered in a trial (the default).
- remove: Unordered timestamps within a trial are removed. This means that any timestamp that is smaller than its predecessor will be removed along with the corresponding x-/y-position.
- ignore: Unordered timestamps are kept and no warning is issued.

\textbf{Value}

A mousetrap data object (see \texttt{mt_example}).

If mouse-tracking data were recorded using the mousetrap plug-ins for OpenSesame, the unit of the timestamps is milliseconds.

\textbf{Author(s)}

Pascal J. Kieslich
Felix Henninger
References


See Also

read_opensesame from the readbulk library for reading and combining raw data files that were collected with OpenSesame.

mt_import_wide and mt_import_long for importing mouse-tracking data from other sources.

Examples

```r
mt_data <- mt_import_mousetrap(mt_example_raw)
```

---

**mt_import_wide**

*Import mouse-tracking data saved in wide format.*

---

Description

mt_import_wide receives a data.frame where mouse-tracking data are stored in wide format, i.e., where one row contains the data of one trial and every recorded mouse position and variable is saved in a separate variable (e.g., X_1, X_2, ..., Y_1, Y_2, ...). This is, e.g., the case when collecting data using MouseTracker (Freeman & Ambady, 2010). From this data.frame, mt_import_wide creates a mousetrap data object containing the trajectories and additional data for further processing within the mousetrap package. Specifically, it returns a list that includes the trajectory data as an array (called trajectories), and all other data as a data.frame (called data). This data structure can then be passed on to other functions within this package (see mousetrap for an overview).

Usage

```r
mt_import_wide(
  raw_data,
  xpos_label = "X",
  ypos_label = "Y",
  zpos_label = NULL,
  timestamps_label = "T",
  add_labels = NULL,
  mt_id_label = NULL,
  pos_sep = ":",
  pos_ids = NULL,
  reset_timestamps = TRUE,
  verbose = TRUE
)
```
Arguments

**raw_data**
a data.frame containing the raw data.

**xpos_label**
a character string specifying the core of the column labels containing the x-positions (e.g., "X" for "X_1", "X_2", ...).

**ypos_label**
a character string specifying the core of the column labels containing the y-positions (e.g., "Y" for "Y_1", "Y_2", ...).

**zpos_label**
a character string specifying the core of the column labels containing the z-positions.

**timestamps_label**
an optional character string specifying the core of the column labels containing the timestamps. If no timestamps are found in the data, a timestamps variable with increasing integers will be created (assuming equidistant time steps).

**add_labels**
a character vector specifying the core of columns containing additional mouse-tracking variables.

**mt_id_label**
an optional character string (or vector) specifying the name of the column that provides a unique ID for every trial (the trial identifier). If unspecified (the default), an ID variable will be generated. If more than one variable name is provided, a new ID variable will be created by combining the values of each variable. The trial identifier will be set as the rownames of the resulting trajectories and trial data, and additionally be stored in the column "mt_id" in the trial data.

**pos_sep**
a character string indicating the character that connects the core label and the position, (e.g., "_" for "X_1", "Y_1", ...).

**pos_ids**
the vector of IDs used for indexing the x-coordinates, y-coordinates etc. (e.g., 1:101 for time-normalized trajectories from MouseTracker). If unspecified (the default), column labels for the respective variable will be extracted using grep (see Details).

**reset_timestamps**
logical indicating if the first timestamp should be subtracted from all timestamps within a trial. Default is `TRUE` as it is recommended for all following analyses in mousetrap.

**verbose**
logical indicating whether function should report its progress.

Details

`mt_import_wide` is designed to import mouse-tracking data saved in a wide format. The defaults are set so that usually only the `raw_data` need to be provided when data have been collecting using MouseTracker (Freeman & Ambady, 2010) and have been read into R using `read_mnt`.

If no pos_ids are provided, column labels for the respective variable (e.g., x-positions) are extracted using `grep` returning all variables that start with the respective character string (e.g., "X_" if xpos_label="X" and pos_sep="_").

If no timestamps are found in the data, `mt_import_wide` automatically creates a timestamps variable with increasing integers (starting with 0) assuming equally spaced sampling intervals.
mt_length_normalize

Value

A mousetrap data object (see mt_example).

Author(s)

Pascal J. Kieslich
Felix Henninger

References


See Also

read_mt for reading raw data that was collected using MouseTracker (Freeman & Ambady, 2010) and stored as a file in the ".mt" format.
mt_import_mousetrap and mt_import_long for importing mouse-tracking data in other formats.

Examples

# Create data in wide format for test purposes
mt_data_wide <- mt_export_wide(mt_example,
   use2_variables=c("subject_nr", "Condition"))

# Import the data using mt_import_wide
mt_data <- mt_import_wide(mt_data_wide,
   xpos_label="xpos", ypos_label="ypos",
   timestamps_label="timestamps")

mt_length_normalize

Length normalize trajectories.

Description

Re-represent each trajectory spatially using a constant number of points so that adjacent points on the trajectory become equidistant to each other.

Usage

mt_length_normalize(
   data,
   use = "trajectories",
   dimensions = c("xpos", "ypos"),
   save_as = "ln_trajectories",
   n_points = 20
)

mt_length_normalize
**Arguments**

- **data**
  a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

- **use**
  a character string specifying which trajectory data should be used.

- **dimensions**
  a character string specifying which trajectory variables should be used. Can be of length 2 or 3 for two-dimensional or three-dimensional data.

- **save_as**
  a character string specifying where the resulting trajectory data should be stored.

- **n_points**
  an integer or vector of integers specifying the number of points used to represent the spatially rescaled trajectories. If a single integer is provided, the number of points will be constant across trajectories. Alternatively, a vector of integers can provided that specify the number of points for each trajectory individually.

**Details**

`mt_length_normalize` is used to emphasize the trajectories' shape. Usually, the vast majority of points of a raw or a time-normalized trajectory lie close to the start and end point. `mt_length_normalize` re-distributes these points so that the spatial distribution is uniform across the entire trajectory. `mt_length_normalize` is mainly used to improve the results of clustering (in particular `mt_cluster`) and visualization.

**Value**

A mousetrap data object (see mt_example) with an additional array (by default called ln_trajectories) containing the length normalized trajectories. If a trajectory array was provided directly as data, only the length normalized trajectories will be returned.

**Author(s)**

Dirk U. Wulff

Jonas M. B. Haslbeck

**Examples**

```r
KH2017 <- mt_length_normalize(data=KH2017, 
  dimensions = c('xpos','ypos'),
  n_points = 20)
```
mt_map

Map trajectories to prototypes.

Description

mt_map maps trajectories onto a predefined set of prototype trajectories. It first computes distances between the trajectories and each of the supplied trajectory types and then assigns each trajectory to the prototype that produced the smallest distance.

Usage

mt_map(
  data,
  use = "ln_trajectories",
  save_as = "prototyping",
  dimensions = c("xpos", "ypos"),
  prototypes = mousetrap::mt_prototypes,
  weights = rep(1, length(dimensions)),
  pointwise = TRUE,
  na_rm = FALSE,
  minkowski_p = 2,
  use2 = "data",
  grouping_variables = NULL
)

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use a character string specifying which trajectory data should be used.

save_as a character string specifying where the resulting data should be stored.

dimensions a character vector specifying which trajectory variables should be used. Can be of length 2 or 3 for two-dimensional or three-dimensional trajectories respectively.

prototypes a trajectory array containing the prototypes the trajectories are mapped to. As a starting point, the trajectories stored in mt_prototypes can be used. See Details and Examples for selecting prototypes and creating new ones.

weights numeric vector specifying the relative importance of the variables specified in dimensions. Defaults to a vector of 1s implying equal importance. Technically, each variable is rescaled so that the standard deviation matches the corresponding value in weights. To use the original variables, set weights = NULL.

pointwise boolean specifying the way dissimilarity between the trajectories is measured (see Details). If TRUE (the default), mt_distmat measures the average dissimilarity and then sums the results. If FALSE, mt_distmat measures dissimilarity once (by treating the various points as independent dimensions).
na.rm logical specifying whether trajectory points containing NAs should be removed. Removal is done column-wise. That is, if any trajectory has a missing value at, e.g., the 10th recorded position, the 10th position is removed for all trajectories. This is necessary to compute distance between trajectories.

minkowski_p an integer specifying the distance metric. minkowski_p = 1 computes the city-block distance, minkowski_p = 2 (the default) computes the Euclidian distance, minkowski_p = 3 the cubic distance, etc.

use2 an optional character string specifying where the data that contain the variables used for grouping can be found (in case grouping_variables are specified). Defaults to "data" as data["data"] usually contains all non mouse-tracking trial data.

grouping_variables a character string (or vector) specifying one or more variables in use2. If specified, prototypes will be rescaled separately to match the coordinate system of the trajectories for each level of the variable(s). If unspecified (the default), the prototypes are rescaled in the same way across all trajectories.

Details

Mouse trajectories often occur in distinct, qualitative types (see Wulff et al., 2019; Wulff et al., 2021). Common trajectory types are linear trajectories, mildly and strongly curved trajectories, and single and multiple change-of-mind trials. mt_map allows to map trajectories to a predefined set of trajectory types.

First, mt_map adjusts prototypes to match the coordinate system of the trajectories specified by use. Next, mt_map computes the distances between each trajectory and each of the supplied prototypes (see mt_distmat) and then assigns each trajectory to the closest prototype (i.e., the prototype that produced the smallest distance).

Mapping trajectories to prototypes requires that the endpoints of all trajectories (and added prototypes) share the same direction, i.e., that all trajectories end in the top-left corner of the coordinate system (mt_remap_symmetric or mt_align can be used to achieve this). Furthermore, it is recommended to use length normalized trajectories (see mt_length_normalize; Wulff et al., 2019).

Value

A mousetrap data object (see mt_example) with an additional data.frame (by default called prototyping) that contains the best fitting prototype for each trajectory (the number of the prototype is specified under prototype, the label of the prototype under prototype_label) and the distance of the trajectory to the best fitting prototype (min_dist). If a trajectory array was provided directly as data, only the data.frame containing the results will be returned.

Author(s)

Dirk U. Wulff
Jonas M. B. Haslbeck
Pascal J. Kieslich
References


Examples

# Length normalize trajectories
KH2017 <- mt_length_normalize(KH2017)

# Map trajectories onto standard prototype set
KH2017 <- mt_map(KH2017,
  use="ln_trajectories")

# Plot prototypes
mt_plot(mt_prototypes,facet_col="mt_id") +
  ggplot2::facet_grid(.~factor(mt_id,levels=unique(mt_id)))

# Plot trajectories per assigned prototype
mt_plot(KH2017,use="ln_trajectories",
  use2="prototyping",facet_col="prototype_label")

# Map trajectories onto reduced prototype set
KH2017 <- mt_map(KH2017,
  use="ln_trajectories",
  prototypes=mt_prototypes[c("straight","curved","cCoM"),,],
  save_as="prototyping_red")

# Map trajectories onto extended prototype set

# Add additional prototypes
mt_prototypes_ext <- mt_add_trajectory(mt_prototypes,
  xpos = c(0,1,-1,1,-1), ypos = c(0,1.5,1.5,1.5), id = "dCoM3")
mt_prototypes_ext <- mt_add_trajectory(mt_prototypes_ext,
  xpos = c(0,0,-1), ypos = c(0,1.5,1.5), id = "neutral")

# Map trajectories
KH2017 <- mt_map(KH2017,
  use="ln_trajectories", prototypes=mt_prototypes_ext,
  save_as="prototyping_ext"
mt_measures

Calculate mouse-tracking measures.

Description

Calculate a number of mouse-tracking measures for each trajectory, such as minima, maxima, and flips for each dimension, and different measures for curvature (e.g., MAD, AD, and AUC). Note that some measures are only returned if distance, velocity and acceleration are calculated using mt_derivatives before running mt_measures. More information on the different measures can be found in the Details and Values sections.

Usage

mt_measures(
  data,
  use = "trajectories",
  save_as = "measures",
  dimensions = c("xpos", "ypos"),
  timestamps = "timestamps",
  flip_threshold = 0,
  hover_threshold = NULL,
  hover_incl_initial = TRUE,
  initiation_threshold = 0,
  verbose = FALSE
)

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use a character string specifying which trajectory data should be used.

save_as a character string specifying where the calculated measures should be stored.

dimensions a character vector specifying the two dimensions in the trajectory array that contain the mouse positions. Usually (and by default), the first value in the vector corresponds to the x-positions (xpos) and the second to the y-positions (ypos).

timestamps a character string specifying the trajectory dimension containing the timestamps.

flip_threshold a numeric value specifying the distance that needs to be exceeded in one direction so that a change in direction counts as a flip. If several thresholds are specified, flips will be returned in separate variables for each threshold value (the variable name will be suffixed with the threshold value).

hover_threshold an optional numeric value. If specified, hovers (and hover_time) will be calculated as the number (and total time) of periods without movement in a trial
(whose duration exceeds the value specified in hover_threshold). If several thresholds are specified, hovers and hover_time will be returned in separate variables for each threshold value (the variable name will be suffixed with the threshold value).

**hover_incl_initial**

logical indicating if the calculation of hovers should include a potential initial phase in the trial without mouse movements (this initial phase is included by default).

**initiation_threshold**

a numeric value specifying the distance from the start point of the trajectory that needs to be exceeded for calculating the initiation time. By default, it is 0, meaning that any movement counts as movement initiation.

**verbose**

logical indicating whether function should report its progress.

### Details

Note that some measures are only returned if distance, velocity and acceleration are calculated using `mt_derivatives` before running `mt_measures`. Besides, the meaning of these measures depends on the values of the arguments in `mt_derivatives`.

If the deviations from the idealized response trajectory have been calculated using `mt_deviations` before running `mt_measures`, the corresponding data in the trajectory array will be used. If not, `mt_measures` will calculate these deviations automatically.

The calculation of most measures can be deduced directly from their definition (see Value). For several more complex measures, a few details are provided in the following.

The signed **maximum absolute deviation** (MAD) is the maximum perpendicular deviation from the straight path connecting start and end point of the trajectory (e.g., Freeman & Ambady, 2010). If the MAD occurs above the direct path, this is denoted by a positive value. If it occurs below the direct path, this is denoted by a negative value. This assumes that the complete movement in the trial was from bottom to top (i.e., the end point has a higher y-position than the start point). In case the movement was from top to bottom, `mt_measures` automatically flips the signs. Both MD_above and MD_below are also reported separately.

The **average deviation** (AD) is the average of all deviations across the trial. Note that AD ignores the timestamps when calculating this average. This implicitly assumes that the time passed between each recording of the mouse is the same within each individual trajectory. If the AD is calculated using raw data that were obtained with an approximately constant logging resolution (sampling rate), this assumption is usually justified (`mt_check_resolution` can be used to check this). Alternatively, the AD can be calculated based on time-normalized trajectories; these can be computed using `mt_time_normalize` which creates equidistant time steps within each trajectory.

The **AUC** represents the **area under curve**, i.e., the geometric area between the actual trajectory and the direct path. Areas above the direct path are added and areas below are subtracted. The AUC is calculated using the `polyarea` function from the pracma package.

Note that all **time** related measures (except idle_time and hover_time) are reported using the timestamp metric as present in the data. To interpret the timestamp values as time since tracking start, the assumption has to be made that for each trajectory the tracking started at timestamp 0 and that all timestamps indicate the time passed since tracking start. Therefore, all timestamps should be reset during data import by subtracting the value of the first timestamp from all timestamps.
within a trial (assuming that the first timestamp corresponds to the time when tracking started). Timestamps are reset by default when importing the data using one of the `mt_import` functions (e.g., `mt_import_mousetrap`). Note that `initiation_time` is defined as the last timestamp before the `initiation_threshold` was crossed.

**Value**

A mousetrap data object (see `mt_example`) where an additional `data.frame` has been added (by default called "measures") containing the per-trial mouse-tracking measures. Each row in the data.frame corresponds to one trajectory (the corresponding trajectory is identified via the row-names and, additionally, in the column "mt_id"). Each column in the data.frame corresponds to one of the measures. If a trajectory array was provided directly as `data`, only the measures data.frame will be returned.

The following measures are computed for each trajectory (the labels relating to x- and y-positions will be adapted depending on the values specified in `dimensions`). Please note that additional information is provided in the Details section.

- **mt_id** Trial ID (can be used for merging measures data.frame with other trial-level data)
- **xpos_max** Maximum x-position
- **xpos_min** Minimum x-position
- **ypos_max** Maximum y-position
- **ypos_min** Minimum y-position
- **MAD** Signed Maximum absolute deviation from the direct path connecting start and end point of the trajectory (straight line). If the MAD occurs above the direct path, this is denoted by a positive value; if it occurs below, by a negative value.
- **MAD_time** Time at which the maximum absolute deviation was reached first
- **MD_above** Maximum deviation above the direct path
- **MD_above_time** Time at which the maximum deviation above was reached first
- **MD_below** Maximum deviation below the direct path
- **MD_below_time** Time at which the maximum deviation below was reached first
- **AD** Average deviation from direct path
- **AUC** Area under curve, the geometric area between the actual trajectory and the direct path where areas below the direct path have been subtracted
- **xpos_flips** Number of directional changes along x-axis (exceeding the distance specified in `flip_threshold`)
- **ypos_flips** Number of directional changes along y-axis (exceeding the distance specified in `flip_threshold`)
- **xpos_reversals** Number of crossings of the y-axis
- **ypos_reversals** Number of crossings of the x-axis
- **RT** Response time, time at which tracking stopped
- **initiation_time** Time at which first mouse movement was initiated
idle_time  Total time without mouse movement across the entirety of the trial
hover_time Total time of all periods without movement in a trial (whose duration exceeds the value specified in hover_threshold)
hovers Number of periods without movement in a trial (whose duration exceeds the value specified in hover_threshold)
total_dist  Total distance covered by the trajectory
vel_max  Maximum velocity
vel_max_time Time at which maximum velocity occurred first
vel_min  Minimum velocity
vel_min_time Time at which minimum velocity occurred first
acc_max  Maximum acceleration
acc_max_time Time at which maximum acceleration occurred first
acc_min  Minimum acceleration
acc_min_time Time at which minimum acceleration occurred first

Author(s)
Pascal J. Kieslich
Felix Henninger

References

See Also
mt_sample_entropy for calculating sample entropy.
mt_standardize for standardizing the measures per subject.
mt_check_bimodality for checking bimodality of the measures using different methods.
mt_aggregate and mt_aggregate_per_subject for aggregating the measures.
inner_join for merging data using the dplyr package.

Examples
mt_example <- mt_derivatives(mt_example)
mt_example <- mt_deviations(mt_example)
mt_example <- mt_measures(mt_example)
# Merge measures with trial data
mt_example_results <- dplyr::inner_join(
mt_example$data, mt_example$measures, by="mt_id")

mt_plot

Plot trajectory data.

Description

mt_plot can be used for plotting a number of individual trajectories. mt_plot_aggregate can be used for plotting aggregated trajectories. The color and linetype can be varied depending on a set of condition variables using the color and linetype arguments. If the x and y arguments are varied, this function can also be used for plotting velocity and acceleration profiles.

Usage

mt_plot(
data,
  use = "trajectories",
  use2 = "data",
  x = "xpos",
  y = "ypos",
  color = NULL,
  linetype = NULL,
  alpha = NA,
  size = 0.5,
  facet_row = NULL,
  facet_col = NULL,
  wrap_var = NULL,
  wrap_ncol = NULL,
  points = FALSE,
  return_type = "plot",
  mt_id = "mt_id",
  only_ggplot = NULL,
  ...
)

mt_plot_aggregate(
data,
  use = "trajectories",
  use2 = "data",
  x = "xpos",
  y = "ypos",
  color = NULL,
  linetype = NULL,
  alpha = NA,
  size = 0.5,
Arguments

- **data**: a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

- **use**: a character string specifying which trajectories should be plotted. The corresponding trajectories are selected from data using data[[use]]. Usually, this value corresponds to either "trajectories", "tn_trajectories" or "av_trajectories", depending on whether the raw, time-normalized or averaged trajectories should be plotted.

- **use2**: a character string specifying where the data that contain the variables used for determining the color and linetype can be found (in case these arguments are specified). Defaults to "data" as data[["data"]] usually contains all non mouse-tracking trial data.

- **x**: a character string specifying which dimension in the trajectory array should be displayed on the x-axis (defaults to xpos).

- **y**: a character string specifying which dimension in the trajectory array should be displayed on the y-axis (defaults to ypos).

- **color**: an optional character string specifying which variable in data[[use2]] should be used for coloring the trajectories.

- **linetype**: an optional character string specifying which variable in data[[use2]] should be used for varying the linetype of the trajectories.

- **alpha**: an optional numeric value between 0 and 1 that can be used to make the plotted lines (and points) semitransparent.

- **size**: an optional numeric value that can be used to vary the width of the plotted trajectory lines.

- **facet_row**: an optional character string specifying a variable in data[[use2]] that should be used for (row-wise) faceting.

- **facet_col**: an optional character string specifying a variable in data[[use2]] that should be used for (column-wise) faceting.

- **wrap_var**: an optional character string specifying variable(s) in data[[use2]] that should be used for wrapping.

- **wrap_ncol**: an optional integer specifying the number of columns if wrapping is used.

- **points**: logical. If TRUE, points will be added to the plot using geom_point.
return_type  a character string specifying which type of object should be returned. If "plot" (the default), a new ggplot is created and the trajectories are plotted using geom_path. If "mapping", only the ggplot object containing the mapping but without any geoms is returned. If "geoms", only the geoms are returned, which allows adding the plotted trajectories to an existing ggplot.

mt_id  a character string specifying the internal label used for the trial identifier (passed on to the group aesthetic). Only relevant for mt_plot.

only_ggplot  Deprecated. Please use return_type instead.

subject_id  a character string specifying which column contains the subject identifier. Only relevant for mt_plot_aggregate. If specified, aggregation will be performed within subjects first. Note that aggregation will be performed separately for each level, including all subjects for whom data are available.

Details

mt_plot internally uses mt_reshape for reshaping trajectories into a long format. Next, it creates a ggplot object using the ggplot function of the ggplot2 package. The aes mappings are taken from the function arguments for x, y etc.; in addition, the group mapping is set to the internal trial identifier (by default called "mt_id").

If return_type == "plot" (the default), a new ggplot is created and the trajectories are plotted using the geom_path function of the ggplot2 package. If return_type == "mapping", the ggplot object is returned without layers, which can be used to further customize the plot (see Examples). If return_type == "geoms", only the geoms are returned, which allows adding the plotted trajectories to an existing ggplot (e.g., adding aggregate trajectories on top of the individual trajectories, see Examples).

mt_plot_aggregate works similarly, but uses mt_aggregate for reshaping and aggregating trajectories prior to plotting.

Please note that this function is intended as a quick and easy solution for visualizing mouse trajectories. For additional flexibility, we recommend that mt_reshape or mt_aggregate be used in conjunction with ggplot to create custom visualizations.

Functions

- mt_plot: Plot individual trajectory data
- mt_plot_aggregate: Plot aggregated trajectory data

Author(s)

Pascal J. Kieslich
Felix Henninger

See Also

mt_plot_add_rect for adding rectangles representing the response buttons to the plot.
m_plot_riverbed for plotting the relative frequency of a selected variable across time.
m_plot_per_trajectory for individually plotting all trajectories as individual pdf files.
Examples

## Plot individual example trajectories

```r
# Time-normalize trajectories
mt_example <- mt_time_normalize(mt_example)

# Plot all time-normalized trajectories
# varying the color depending on the condition
mt_plot(mt_example, use="tn_trajectories",
       color="Condition")

# ... setting alpha < 1 for semi-transparency
mt_plot(mt_example, use="tn_trajectories",
       color="Condition", alpha=.2)

# ... with custom colors
mt_plot(mt_example, use="tn_trajectories",
       color="Condition") +
  ggplot2::scale_color_brewer(type="qual")

# Create separate plots per Condition
mt_plot(mt_example, use="tn_trajectories",
       facet_col="Condition")

# Create customized plot by setting the return_type option to "mapping"
# to setup an empty plot. In a next step, a geom is added.
# In this example, only points are plotted.
# Plot velocity profiles based on the averaged trajectories
# varying the color depending on the condition
mt_example <- mt_derivatives(mt_example)
mt_example <- mt_average(mt_example, interval_size=100)
mt_plot(mt_example, use="av_trajectories",
        x="timestamps", y="vel", color="Condition")
```

## Plot aggregate trajectories for KH2017 data

```r
# Time-normalize trajectories
KH2017 <- mt_time_normalize(KH2017)

# Plot aggregated time-normalized trajectories per condition
mt_plot_aggregate(KH2017, use="tn_trajectories",
                 color="Condition")

# ... first aggregating trajectories within subjects
mt_plot_aggregate(KH2017, use="tn_trajectories",
                 color="Condition", subject_id="subject_nr")
```
# ... adding points for each position to the plot
mt_plot_aggregate(KH2017, use="tn_trajectories",
    color="Condition", points=TRUE)

## Not run:

# Create combined plot of individual and aggregate trajectories
# by first plotting the individual trajectories using mt_plot.
# In a next step, the aggregate trajectories are added using the
# mt_plot_aggregate function with the return_type argument set to "geom".
mt_plot(KH2017, use="tn_trajectories", color="Condition", alpha=.05) +
  mt_plot_aggregate(KH2017, use="tn_trajectories",
    color="Condition", return_type="geom", size=2)

## End(Not run)

---

### mt_plot_add_rect

**Add rectangles to trajectory plot.**

**Description**

`mt_plot_add_rect` adds one or several rectangles to a mousetrap plot. These buttons usually correspond to the borders of the buttons in the mouse-tracking experiment. It is specifically designed so that the arguments from the `mousetrap_response` plugin in OpenSesame can be used.

**Usage**

```r
mt_plot_add_rect(rect, color = "black", fill = NA, ...)
```

**Arguments**

- `rect`: a data.frame or matrix with one row per box. For each rectangle, the x-position (x), y-position (y), width (w), and height (h) needs to be provided. If columns are not labeled, the order x, y, w, h is assumed.
- `color`: argument passed on to `geom_rect`. Specifies the color of the border of the rectangles.
- `fill`: argument passed on to `geom_rect`. Specifies the color of the interior of the rectangles. If NA (the default), rectangles are unfilled.
- `...`: additional arguments passed on to `geom_rect`.

**Details**

`mt_plot_add_rect` internally uses `geom_rect` of the `ggplot2` package for plotting.

**Author(s)**

Pascal J. Kieslich
Felix Henninger
See Also

`mt_plot` for plotting trajectory data.

Examples

```r
# Load ggplot2
library(ggplot2)

# Import, flip, and time-normalize raw trajectories
mt_example <- mt_import_mousetrap(mt_example_raw)
mt_example <- mt_remap_symmetric(mt_example, remap_xpos="no")
mt_example <- mt_time_normalize(mt_example)

# Create rectangles matrix
rectangles <- matrix(
  c(-840, 525, 350, -170,
    840, 525, -350, -170),
  ncol=4, byrow=TRUE)

# Plot all time-normalized trajectories
# varying the color depending on the condition
# and add rectangles
mt_plot(mt_example,
  use="trajectories",
  x="xpos", y="ypos", color="Condition"
) + mt_plot_add_rect(rect=rectangles)
```

`mt_plot_per_trajectory`

Create PDF with separate plots per trajectory.

Description

`mt_plot_per_trajectory` creates a PDF file with separate plots per trajectory. This PDF can be used for inspecting individual trajectories. Note that plotting all trajectories can be time-consuming, especially for raw trajectories. If the appropriate x and y arguments are inserted, this function can also be used for plotting velocity and acceleration profiles.
mt_plot_per_trajectory

Usage

mt_plot_per_trajectory(
  file,
  data,
  use = "trajectories",
  x = "xpos",
  y = "ypos",
  xlim = NULL,
  ylim = NULL,
  axes_exact = FALSE,
  points = FALSE,
  rect = NULL,
  color = "black",
  fill = NA,
  verbose = FALSE,
  ...
)

Arguments

file a character string specifying the name of the PDF file. Passed on to pdf.
data a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).
use a character string specifying which trajectories should be plotted. The corresponding trajectories are selected from data using `data[[use]]`. Usually, this value corresponds to either "trajectories", "tn_trajectories" or "av_trajectories", depending on whether the raw, time-normalized or averaged trajectories should be plotted.
x a character string specifying which dimension in the trajectory array should be displayed on the x-axis (defaults to xpos).
y a character string specifying which dimension in the trajectory array should be displayed on the y-axis (defaults to ypos).
xlim optional argument specifying the limits for the x axis (passed on to coord_cartesian). If not specified (the default), sensible axis limits will be computed.
ylim optional argument specifying the limits for the y axis (passed on to coord_cartesian). If not specified (the default), sensible axis limits will be computed.
axes_exact logical. If TRUE, axes will be set without offset exactly at the limits of the x and y axes (which can be specified using xlim and ylim).
points logical. If TRUE, points will be added to the plot using geom_point.
rect optional argument passed on to mt_plot_add_rect. If specified, rectangles (usually representing the response buttons) will be plotted for each trajectory plot.
color optional argument passed on to mt_plot_add_rect. Only relevant if rect is specified.
mt_plot_riverbed  

 Plot density of mouse positions across time steps.

Description

mt_plot_riverbed creates a plot showing the distribution of one trajectory variable (e.g., the x-positions or velocity) per time step.

Usage

mt_plot_riverbed(  
data,  
use = "tn_trajectories",  
y = "xpos",  
y_range = NULL,  
y_bins = 250,  
facet_row = NULL,  
facet_col = NULL,
mt_plot_riverbed

```r
facenet_data = "data",
grid_colors = c("gray30", "gray10"),
na.rm = FALSE
)
```

Arguments

- **data**: mousetrap data object containing the data to be plotted.
- **use**: character string specifying the set of trajectories to use in the plot. The steps of this set will constitute the x axis. Defaults to ‘tn_trajectories’, which results in time steps being plotted on the x axis.
- **y**: variable in the mousetrap data object to be plotted on the output’s y dimension. Defaults to ‘xpos’, the cursor’s x coordinate.
- **y_range**: numerical vector containing two values that represent the upper and lower ends of the y axis. By default, the range is calculated from the data provided.
- **y_bins**: number of bins to distribute along the y axis (defaults to 250).
- **facet_row**: an optional character string specifying a variable in data[[facet_data]] that should be used for (row-wise) faceting. If specified, separate riverbed plots for each level of the variable will be created.
- **facet_col**: an optional character string specifying a variable in data[[facet_data]] that should be used for (column-wise) faceting. If specified, separate riverbed plots for each level of the variable will be created.
- **facet_data**: a character string specifying where the (optional) data containing the faceting variables can be found.
- **grid_colors**: a character string or vector of length 2 specifying the grid color(s). If a single value is provided, this will be used as the grid color. If a vector of length 2 is provided, the first value will be used as the color for the major grid lines, the second value for the minor grid lines. If set to NA, no grid lines are plotted.
- **na.rm**: logical specifying whether missing values should be removed. This is not done by default, because generally riverbed plots are generated from preprocess trajectories (e.g., time-normalized trajectories) that all have the same length (i.e., the same number of steps).

Details

This function plots the relative frequency of the values of a trajectory variable separately for each of a series of time steps. This type of plot has been used in previous research to visualize the distribution of x-positions per time step (e.g., Scherbaum et al., 2010).

mt_plot_riverbed usually is applied to time-normalized trajectory data as all trajectories must contain the same number of values (if na.rm=FALSE, the default).

Author(s)

Felix Henninger
Pascal J. Kieslich
References


See Also

- `mt_plot` for plotting trajectory data.
- `mt_time_normalize` for time-normalizing trajectories.

Examples

```r
# Time-normalize trajectories
KH2017 <- mt_time_normalize(KH2017)

# Create riverbed plot for all trials
mt_plot_riverbed(KH2017)

## Not run:
# Create separate plots for typical and atypical trials
mt_plot_riverbed(mt_example, facet_col="Condition")

# Create riverbed plot for all trials with custom x and y axis labels
mt_plot_riverbed(mt_example) +
  ggplot2::xlab("Time step") + ggplot2::ylab("X coordinate")

# Note that it is also possible to replace the
# default scale for fill with a custom scale
mt_plot_riverbed(mt_example, facet_col="Condition") +
  ggplot2::scale_fill_gradientn(colours=grDevices::heat.colors(9),
                              name="Frequency", trans="log", labels=scales::percent)

## End(Not run)
```

---

**mt_prototypes**

*Mouse trajectory prototypes.*

Description

A core set of five mouse trajectory prototypes including the ‘straight’ trajectory, the mildly curved trajectory, the continuous change-of-mind trajectory, the discrete change-of-mind trajectory, and the double discrete change-of-mind trajectory.

Usage

`mt_prototypes`
Format

An object of class array of dimension 5 x 100 x 2.

Details

Mouse- and hand-trajectories often occur in types (Wulff, Haslbeck, & Schulte-Mecklenbeck, 2017). In such cases, movement trajectory data should be analyzed in terms of discrete type assignments. To this end mt_map can be used to map mouse- or hand-trajectory to the closest of several predefined prototypes. mt_prototypes provides a core set of prototypes that has been shown to represent well a large fraction of empirical movement trajectories.

To tailor the set of prototypes to a given study, mt_prototypes can be extended using mt_add_trajectory.

References


---

mt_qeffect Create quantile-effect plot

Description

[Experimental]

Function in beta and currently only for internal purposes.

Usage

```r
mt_qeffect(
  data,
  compare,
  use = "measures",
  measure = "MAD",
  direction = "upward",
  n_steps = 100,
  return_data = FALSE,
  ...
)
```

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

compare either a vector, the label of a variable in , or a mousetrap object.

use a character string specifying which trajectory data should be used.
**mt_remap_symmetric**

Remap mouse trajectories.

Description

Remap all trajectories to one side (or one quadrant) of the coordinate system. In doing so, *mt_remap_symmetric* assumes a centered coordinate system and a symmetric design of the response buttons (see Details).

Usage

```r
mt_remap_symmetric(
  data,
  use = "trajectories",
  save_as = use,
  dimensions = c("xpos", "ypos"),
  remap_xpos = "left",
  remap_ypos = "up"
)
```

**Value**

Nothing, when image is plotted using an external device. Otherwise an object of class `mt_object_raw` containing in a matrix format the image’s pixel information.

**Author(s)**

Dirk U. Wulff

**Examples**

```r
# Plot regular heatmap
#SpiveyEtAl2005 = mt_import_long(SpiveyEtAl2005_raw, 'x', 'y', NULL, 't',
#mt_id_label = c('ptp', 'trial'))
#heatmap = mt_heatmap_raw(SpiveyEtAl2005, xres = 2000)
#mt_heatmap(heatmap, file = NULL)

# compute measures
#SpiveyEtAl2005 = mt_measures(SpiveyEtAl2005)

# Plot heatmap using velocity
#mt_heatmap(SpiveyEtAl2005)
```
Arguments

**data**  
a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

**use**  
a character string specifying which trajectory data should be used.

**save_as**  
a character string specifying where the resulting trajectory data should be stored.

**dimensions**  
a character vector specifying the two dimensions in the trajectory array that contain the mouse positions, the first value corresponding to the x-positions, the second to the y-positions.

**remap_xpos**  
character string indicating the direction in which to remap values on the x axis. If set to "left" (as per default), trajectories with an endpoint on the right (i.e. with a positive x-value) will be remapped to the left. The alternatives are "right" which has the reverse effect, and "no", which disables remapping on the horizontal dimension.

**remap_ypos**  
character string defining whether tracks directed downwards on the y axis should be remapped so that they end with a positive y value. This will be performed if this parameter is set to "up" (which is the default), and the reverse occurs if the parameter is set to "down". If it is set to "no", y-values remain untouched.

Details

When mouse trajectories are compared across different conditions, it is typically desirable that the endpoints of the trajectories share the same direction (e.g., diagonally up and left). This way, the trajectories can be compared regardless of the button they were directed at.

**mt_remap_symmetric** can be used to achieve this provided that two assumptions hold:

First, this function assumes a centered coordinate system, i.e. the coordinate system is centered on the screen center. This is the case when the data is produced by the mousetrap plug-ins in OpenSesame.

Second, it assumes that the response buttons in the mouse-tracking experiment are symmetric, in that they all are equally distant from the screen center.

Value

A mousetrap data object (see mt_example) with remapped trajectories. If the trajectory array was provided directly as data, only the trajectory array will be returned.

Author(s)

Pascal J. Kieslich
Felix Henninger

Examples

# Remap trajectories so that all trajectories  
# end in the top-left corner  
mt_example <- mt_import_mousetrap(mt_example_raw)
\begin{verbatim}
mt_example <- mt_remap_symmetric(mt_example)

# Only flip trajectories vertically so that all
# trajectories end in the upper half of the screen
mt_example <- mt_import_mousetrap(mt_example_raw)
mt_example <- mt_remap_symmetric(mt_example,
    remap_xpos="no", remap_ypos="up")
\end{verbatim}

\textbf{mt_resample} \hspace{1cm} \textit{Resample trajectories using a constant time interval.}

\textbf{Description}

Resample trajectory positions using a constant time interval. If no timestamp that represents an exact multiple of this time interval is found, linear interpolation is performed using the two adjacent timestamps.

\textbf{Usage}

\begin{verbatim}
mt_resample(  
data,  
    use = "trajectories",  
    save_as = "rs_trajectories",  
    dimensions = c("xpos", "ypos"),  
    timestamps = "timestamps",  
    step_size = 10,  
    exact_last_timestamp = TRUE,  
    constant_interpolation = NULL,  
    verbose = FALSE
)
\end{verbatim}

\textbf{Arguments}

\begin{itemize}
  \item \textbf{data} \hspace{1cm} a mousetrap data object created using one of the \texttt{mt_import} functions (see \texttt{mt_example} for details). Alternatively, a trajectory array can be provided directly (in this case \texttt{use} will be ignored).
  \item \textbf{use} \hspace{1cm} a character string specifying which trajectory data should be used.
  \item \textbf{save_as} \hspace{1cm} a character string specifying where the resulting trajectory data should be stored.
  \item \textbf{dimensions} \hspace{1cm} a character vector specifying the dimensions in the trajectory array that should be resampled. If "all", all trajectory dimensions except the timestamps will be resampled.
  \item \textbf{timestamps} \hspace{1cm} a character string specifying the trajectory dimension containing the timestamps.
  \item \textbf{step_size} \hspace{1cm} an integer specifying the size of the constant time interval. The unit corresponds to the unit of the timestamps.
\end{itemize}
exact_last_timestamp

logical indicating if the last timestamp should always be appended (which is the case by default). If FALSE, the last timestamp is only appended if it is a multiple of the step_size.

constant_interpolation

an optional integer. If specified, constant instead of linear interpolation will be performed for all adjacent timestamps whose difference exceeds the number specified for constant_interpolation. The unit corresponds to the unit of the timestamps.

verbose

logical indicating whether function should report its progress.

Details

mt_resample can be used if the number of logged positions in a trial should be reduced. mt_resample achieves this by artificially decreasing the resolution with which the positions were recorded. For example, if mouse positions were recorded every 10 ms in an experiment, but one was only interested in the exact mouse position every 50 ms, mt_resample with step_size=50 could be used. In this case, only every fifth sample would be kept.

In addition, mt_resample can be used to only retain values for specific timestamps across trials (e.g., if for each trial the position of the mouse exactly 250 ms and 500 ms after onset of the trial are of interest). In case that a trial does not contain samples at the specified timestamps, linear interpolation is performed using the two adjacent timestamps.

If a number is specified for constant_interpolation, constant instead of linear interpolation will be performed for all adjacent timestamps whose difference exceeds this number. Specifically, a period without mouse movement will be assumed starting at the respective timestamp until the next timestamp

• constant_interpolation/2.

Note that mt_resample does not average across time intervals. For this, mt_average can be used.

Value

A mousetrap data object (see mt_example) with an additional array (by default called rs_trajectories) containing the resampled trajectories. If a trajectory array was provided directly as data, only the resampled trajectories will be returned.

Author(s)

Pascal J. Kieslich
Felix Henninger

See Also

approx for information about the function used for linear interpolation.
mt_average for averaging trajectories across constant time intervals.
mt_time_normalize for time-normalizing trajectories.
Examples

```r
mt_example <- mt_resample(mt_example,
save_as="rs_trajectories",
step_size=50)
```

### Description

`mt_reshape` is the general function used in the mousetrap package for filtering, merging, reshaping, and aggregating mouse-tracking measures or trajectories in combination with other trial data. Several additional (wrapper) functions for more specific purposes (cf. "See Also") are available.

### Usage

```r
mt_reshape(
  data,
  use = "trajectories",
  use_variables = NULL,
  use2 = "data",
  use2_variables = NULL,
  subset = NULL,
  subject_id = NULL,
  aggregate = FALSE,
  aggregate_subjects_only = FALSE,
  .funs = "mean",
  trajectories_long = TRUE,
  convert_df = TRUE,
  mt_id = "mt_id",
  mt_seq = "mt_seq",
  aggregation_function = NULL
)
```

### Arguments

- **data**: a mousetrap data object created using one of the mt_import functions (see `mt_example` for details). Alternatively, a trajectory array can be provided directly (in this case `use` will be ignored).
- **use**: a character string specifying which data should be reshaped. The corresponding data are selected from `data` using `data[[use]]`. Usually, this value corresponds to either "trajectories", "tn_trajectories", or "measures", depending on whether the analysis concerns raw trajectories, time-normalized trajectories, or derived measures.
use_variables  a character vector specifying which mouse-tracking variables should be reshaped. Corresponds to the column names in case a data.frame with mouse-tracking measures is provided. Corresponds to the labels of the array dimensions in case a trajectory array is provided. If unspecified, all variables will be reshaped.

use2  an optional character string specifying where the other trial data can be found. Defaults to "data" as data[["data"]]) usually contains all non mouse-tracking trial data. Alternatively, a data.frame can be provided directly.

use2_variables  an optional character string (or vector) specifying the variables (in data[[use2]]) that should be merged with the data. If aggregate==TRUE, the trajectories/measures will be aggregated separately for each of the levels of these variables using summarize_at.

subset  a logical expression (passed on to subset) indicating elements or rows to keep. If specified, data[[use2]] will be subsetted using this expression, and, afterwards, data[[use]] will be filtered accordingly.

subject_id  an optional character string specifying which column contains the subject identifier in data[[use2]]. If specified and aggregate==TRUE, aggregation will be performed within subjects first.

aggregate  logical indicating whether data should be aggregated. If use2_variables are specified, aggregation will be performed separately for each of the levels of the use2_variables.

aggregateSubjectsOnly  logical indicating whether data should only be aggregated per subject (if subject_id is specified and aggregate==TRUE).

.funs  the aggregation function(s) passed on to summarize_at. By default, the mean is calculated.

trajectories_long  logical indicating if the reshaped trajectories should be returned in long or wide format. If TRUE, every recorded position in a trajectory is placed in another row (whereby the order of the positions is logged in the variable mt_seq). If FALSE, every trajectory is saved in wide format and the respective positions are indexed by adding an integer to the corresponding label (e.g., xpos_1, xpos_2,...). Only relevant if data[[use]] contains trajectories.

convert_df  logical indicating if the reshaped data should be converted to a data.frame using as.data.frame. This will drop potentially existing additional classes (such as tbl_df) that result from the internally used dplyr functions for data grouping and aggregation. As these additional classes might cause problems with functions from other packages, the reshaped data are converted to "pure" data.frames by default.

mt_id  a character string specifying the name of the column that will contain the trial identifier in the reshaped data. The values for the trial identifier correspond to the rownames of data[[use]] and data[[use2]].

mt_seq  a character string specifying the name of the column that will contain the integers indicating the order of the mouse positions per trajectory in the reshaped data. Only relevant if data[[use]] contains trajectories and trajectories_long==TRUE.

aggregation_function  Deprecated. Please use .funs instead.
**Details**

`mt_reshape` uses the `rownames` of `data[[use]]` and `data[[use2]]` for merging the trajectories / measures and the trial data. For convenience (and for trajectories in long format also of necessity), an additional column (labelled as specified in the `mt_id` argument) is added to the reshaped data containing the rownames as trial identifier.

The main purpose of this function is to reshape the trajectory data into a two-dimensional data.frame, as this format is required for many further analyses and plots in R.

Besides, it should aid the user in combining data contained in different parts of the mousetrap data object, e.g., a condition variable stored in `data["data"]` with trajectory data stored in `data["trajectories"]` (or mouse-tracking measures stored in `data["measures"]`).

Finally, it offers the possibility to aggregate trajectories and measures for different conditions and/or subjects.

The package also includes several functions that wrap `mt_reshape` and serve specific purposes. They are often easier to use, and thus recommended over `mt_reshape` unless the utmost flexibility is required. These functions are described in the section "See Also".

Note also that many merging, reshaping, and aggregation procedures can be performed directly by using some of the basic R functions, e.g., `merge` and `aggregate`, or through the R packages `dplyr` or `reshape2`, if desired.

**Value**

A `data.frame` containing the reshaped data.

**Author(s)**

Pascal J. Kieslich
Felix Henninger

**See Also**

`mt_aggregate` for aggregating mouse-tracking measures and trajectories.
`mt_aggregate_per_subject` for aggregating mouse-tracking measures and trajectories per subject.
`mt_export_long` for exporting mouse-tracking data in long format.
`mt_export_wide` for exporting mouse-tracking data in wide format.
`inner_join` for merging data and `summarize_at` for aggregating data using the `dplyr` package.

**Examples**

```r
# Time-normalize trajectories
mt_example <- mt_time_normalize(mt_example)

# Reshape time-normalized trajectories data into long format
# adding Condition variable
trajectories_long <- mt_reshape(mt_example, 
    use="tn_trajectories", 
    use2_variables="Condition"
```
mt_sample_entropy

Calculate sample entropy.

Description

Calculate sample entropy for each trajectory as a measure of the complexity of movements along one specific dimension.

Usage

mt_sample_entropy(
  data,
  use = "tn_trajectories",
  save_as = "measures",
  dimension = "xpos",
  m = 3,
  r = NULL,
  use_diff = TRUE,
  verbose = FALSE
)

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use a character string specifying which trajectory data should be used.

save_as a character string specifying where the calculated measures should be stored.

dimension a character string specifying the dimension based on which sample entropy should be calculated. By default (xpos), the x-positions are used.

m an integer passed on to the sample entropy function (see Details).

r a numeric value passed on to the sample entropy function (see Details).

use_diff logical indicating if the differences of the dimension values should be computed before calculating sample entropy (which is done by default, see Details).

verbose logical indicating whether function should report its progress.
**Details**

`mt_sample_entropy` calculates the sample entropy for each trajectory as a measure of its complexity. Hehman et al (2015) provide details on how sample entropy can be calculated and applied in mouse-tracking (following Dale et al., 2007). They apply the sample entropy measure to the differences between adjacent x-positions (which is also the default here, as in a standard mouse-tracking task with buttons located in the top-left and right corners mostly the movements in the horizontal direction are of interest). Besides, they recommend using the time-normalized trajectories so all trajectories have the same length.

Sample entropy is computed by comparing windows of a fixed size (specified using `m`) across all recorded positions. Sample entropy is the negative natural logarithm of the conditional probability that this window remains similar across the trial (Hehman et al., 2015). A window is considered to be similar to another if their distance is smaller than a specified tolerance value (which can be specified using `r`). Hehman et al. (2015) use a tolerance value of 0.2 * standard deviation of all differences between adjacent x-positions in the dataset (which is the default implemented here).

**Value**

A mousetrap data object (see `mt_example`).

If a data.frame with label specified in `save_as` (by default "measures") already exists, the sample entropy values are added as additional column.

If not, an additional data.frame will be added.

If a trajectory array was provided directly as data, only the data.frame will be returned.

**Author(s)**

Pascal J. Kieslich
Dirk Wulff
Felix Henninger

**References**


**See Also**

`mt_measures` for calculating other mouse-tracking measures.

**Examples**

```r
# Calculate sample entropy based on time-normalized
# trajectories and merge results with other measures
# derived from raw trajectories
mt_example <- mt_measures(mt_example)
mt_example <- mt_time_normalize(mt_example,
```

mt_scale_trajectories

Standardize variables in mouse trajectory array.

Description

mt_scale_trajectories centers and / or standardizes selected trajectory variables within or across trajectories.

Usage

mt_scale_trajectories(
  data,
  use = "trajectories",
  save_as = use,
  var_names,
  center = TRUE,
  scale = TRUE,
  within_trajectory = FALSE,
  prefix = "z_",
  transform = NULL
)

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use a character string specifying which trajectory data should be used.

save_as a character string specifying where the resulting trajectory data should be stored.

var_names character vector giving the labels of the to be standardized variables.

center logical specifying whether variables should be centered (i.e., mean = 0). Can be a logical vector, in which case the values of scale are mapped to the variables specified in var_names.

scale logical or numeric specifying the scaling of the variables. When logical, scale = TRUE normalizes the trajectory variable to sd = 1, whereas scale = FALSE leaves the variable on its original scale. When numeric, the trajectory variables are scaled by (i.e., divided by) the specific value in scale. Can also be a numeric vector, in which case the values of scale are mapped to the variables specified in var_names.
within_trajectory

logical specifying whether trajectory variables should be scaled within or across trajectories. If `within_trajectory == TRUE`, scaling trajectories to mean = 0 and sd = 1 means that every to be standardized trajectory variable will have mean = 0 and sd = 1. If `within_trajectory == FALSE` (the default), mean = 0 and sd = 1 are only true in the aggregate (i.e., across all trajectories). Can be a logical vector, in which case the values of `scale` are mapped to the variables specified in `var_names`.

prefix

character string added to the names of the new standardized variables. If `prefix = ""`, the original variables will be overwritten.

transform

function that takes a numeric matrix as argument and returns a numeric matrix of same size with transformed values. If `NULL` the original values are passed on to standardization.

Value

A mousetrap data object (see `mt_example`) with an additional variable containing the standardized trajectory variable added to the trajectory array). If the trajectory array was provided directly as data, only the trajectory array will be returned.

Author(s)

Dirk U. Wulff

See Also

`mt_standardize` for standardizing mouse-tracking measures per level of other variables.

Examples

```r
# Calculate derivatives
mt_example <- mt_derivatives(mt_example)

# Standardize velocity across trajectories
mt_example <- mt_scale_trajectories(mt_example, var_names = "vel")
```

Description

[Deprecated]

Re-represent each trajectory spatially using a constant number of points so that adjacent points on the trajectory become equidistant to each other. Please note that this function is deprecated and that `mt_length_normalize` should be used instead.
Usage

```r
mt_spatialize(
  data,
  use = "trajectories",
  dimensions = c("xpos", "ypos"),
  save_as = "sp_trajectories",
  n_points = 20
)
```

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use a character string specifying which trajectory data should be used.

dimensions a character string specifying which trajectory variables should be used. Can be of length 2 or 3 for two-dimensional or three-dimensional data.

save_as a character string specifying where the resulting trajectory data should be stored.

n_points an integer or vector of integers specifying the number of points used to represent the spatially rescaled trajectories. If a single integer is provided, the number of points will be constant across trajectories. Alternatively, a vector of integers can provided that specify the number of points for each trajectory individually.

Details

`mt_spatialize` is used to emphasize the trajectories’ shape. Usually, the vast majority of points of a raw or a time-normalized trajectory lie close to the start and end point. `mt_spatialize` re-distributes these points so that the spatial distribution is uniform across the entire trajectory. `mt_spatialize` is mainly used to improve the results of clustering (in particular `mt_cluster`) and visualization.

Value

A mousetrap data object (see `mt_example`) with an additional array containing the spatialized trajectories. If a trajectory array was provided directly as data, only the spatialized trajectories will be returned.

Author(s)

Dirk U. Wulff
Jonas M. B. Haslbeck

Examples

```r
## Not run:
KH2017 <- mt_spatialize(data=KH2017,
  dimensions = c("xpos","ypos"),
  n_points = 20)
```
mt_standardize

Standardize selected mouse-tracking measures across all trials or per level of one or more other variable, and store them in new variables. This function is a thin wrapper around scale_within, focussed on mouse-tracking data stored in a mousetrap data object.

Usage

mt_standardize(data, 
use = "measures", 
use_variables = NULL, 
within = NULL, 
prefix = "z_", 
center = TRUE, 
scale = TRUE)

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details).
use a character string specifying which data should be used. By default points to the measures data.frame created using mt_measures.
use_variables a vector specifying which variables should be standardized. If unspecified, all variables will be standardized.
within an optional character string specifying one or more variables in data[["data"]]. If specified, all measures will be standardized separately for each level of the variable (or for each combination of levels, if more than one variable is specified).
prefix a character string that is inserted before each standardized variable. If an empty string is specified, the original variables are replaced.
center argument passed on to scale.
scale argument passed on to scale.

Value

A mousetrap data object (see mt_example) including the standardized measures.
Author(s)

Pascal J. Kieslich
Felix Henninger

See Also

mt_scale_trajectories for standardizing variables in mouse trajectory arrays.
scale_within which is called by mt_standardize.
scale for the R base scale function.

Examples

mt_example <- mt_measures(mt_example)

# Standardize MAD and AD per subject
mt_example <- mt_standardize(mt_example,
   use_variables=c("MAD", "AD"),
   within="subject_nr", prefix="z_")

# Standardize MAD and AD per subject and Condition
mt_example <- mt_standardize(mt_example,
   use_variables=c("MAD", "AD"),
   within=c("subject_nr", "Condition"),
   prefix="z_")

mt_subset

Filter mousetrap data.

Description

Return a subset of the mousetrap data including only the trial data and corresponding trajectories that meet the conditions specified in the arguments.

Usage

mt_subset(data, subset, check = "data")

Arguments

data a mousetrap data object created using one of the mt_import functions (see mt_example for details).

subset a logical expression (passed on to subset) indicating the rows to keep. Missing values are taken as FALSE.

check a character string specifying which data should be used for checking the subset condition.
Details

`mt_subset` is helpful when trials should be removed from all analyses. By default, check is set to "data" meaning that the subset condition is evaluated based on the trial data (stored in `data[["data"]])`. However, it might also be of interest to only include trials based on specific mouse-tracking measures (e.g., all trials with an MAD smaller than 200). In this case, check needs to be set to the respective name of the data.frame (e.g., "measures").

Note that if specific trials should be removed from all analyses based on a condition known a priori (e.g., practice trials), it is more efficient to use the `subset` function on the raw data before importing the trajectories using one of the `mt_import` functions (such as `mt_import_mousetrap`).

Besides, if trials should only be removed from some analyses or for specific plots, note that other mousetrap functions (e.g., `mt_reshape`, `mt_aggregate`, and `mt_plot`) also allow for subsetting.

Value

A mousetrap data object (see `mt_example`) with filtered data and trajectories.

Author(s)

Pascal J. Kieslich
Felix Henninger

See Also

`subset` for the R base subset function for vectors, matrices, or data.frames.

`mt_reshape` for information about the subset argument in various other mousetrap functions.

Examples

```r
# Subset based on trial data
mt_example_atypical <- mt_subset(mt_example, Condition="Atypical")

# Subset based on mouse-tracking measure (MAD)
mt_example <- mt_measures(mt_example)
mt_example_mad_sub <- mt_subset(mt_example, MAD<400, check="measures")
```

Description

Compute time-normalized trajectories using a constant number of equally sized time steps. Time normalization is performed separately for all specified trajectory dimensions (by default, the x- and y-positions) using linear interpolation based on the timestamps. By default, 101 time steps are used (following Spivey et al., 2005).
Usage

mt_time_normalize(
  data,
  use = "trajectories",
  save_as = "tn_trajectories",
  dimensions = c("xpos", "ypos"),
  timestamps = "timestamps",
  nsteps = 101,
  verbose = FALSE
)

Arguments

data             a mousetrap data object created using one of the mt_import functions (see mt_example for details). Alternatively, a trajectory array can be provided directly (in this case use will be ignored).

use             a character string specifying which trajectory data should be used.

save_as         a character string specifying where the resulting trajectory data should be stored.

dimensions       a character vector specifying the dimensions in the trajectory array that should be time-normalized. If "all", all trajectory dimensions except the timestamps will be time-normalized.

timestamps       a character string specifying the trajectory dimension containing the timestamps.

nsteps           an integer specifying the number of equally sized time steps.

verbose         logical indicating whether function should report its progress.

Details

Time-normalization is often performed if the number of recorded x- and y-positions varies across trajectories, which typically occurs when trajectories vary in their response time. After time-normalization, all trajectories have the same number of recorded positions (which is specified using nsteps) and the positions at different relative time points can be compared across trajectories.

For example, time normalized trajectories can be compared across conditions that differed in their overall response time, as the timestamps are now relative to the overall trial duration. This is also helpful for creating average trajectories, which are often used in plots.

Value

A mousetrap data object (see mt_example) with an additional array (by default called tn_trajectories) containing the time-normalized trajectories. In this array, another dimension (called steps) has been added with increasing integer values indexing the time-normalized position. If a trajectory array was provided directly as data, only the time-normalized trajectories will be returned.

Author(s)

Pascal J. Kieslich
Felix Henninger
References

See Also
approx for information about the function used for linear interpolation.
mt_resample for resampling trajectories using a constant time interval.

Examples
```
mt_example <- mt_time_normalize(mt_example,
    save_as="tn_trajectories", nsteps=101)
```

print.mt_heatmap_raw

Generic print for class mt_heatmap_raw

Description
print.mt_heatmap_raw shows str.

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

Arguments

<table>
<thead>
<tr>
<th>x</th>
<th>an object of class mt_heatmap_raw.</th>
</tr>
</thead>
<tbody>
<tr>
<td>...</td>
<td>further arguments passed to or from other methods.</td>
</tr>
</tbody>
</table>

read_mt

*Read MouseTracker raw data.*

Description
read_mt reads raw data that was collected using MouseTracker (Freeman & Ambady, 2010) and stored as a file in the ".mt" format. If multiple files should be read into R, read_mt can be used in combination with the read_bulk function from the readbulk package (see Examples). After reading the data into R, mt_import_wide can be used to prepare the trajectory data for analyses using the mousetrap library. The current version of read_mt has been tested with data from MouseTracker Version 2.84 - but please be sure to double-check.
Usage

read_mt(file, columns = "all", add_trialid = FALSE, add_filename = FALSE)

Arguments

- **file**: a character string specifying the filename of the .mt file.
- **columns**: either 'all' or a character vector specifying the to be extracted variables. Defaults to 'all' in which case all existing variables will be extracted.
- **add_trialid**: boolean specifying whether an additional column containing the trial number should be added.
- **add_filename**: boolean specifying whether an additional column containing the file name should be added.

Value

A data.frame with one row per trial. Variables are ordered according to columns, x-coordinates, y-coordinates, and timestamps.

Author(s)

Dirk U. Wulff

References


See Also

- `read_bulk` from the readbulk package for reading and combining multiple raw data files.
- `mt_import_wide` to prepare the imported data for analyses in mousetrap.

Examples

```r
## Not run:
# Read a single raw data file from MouseTracker
# (stored in the current working directory)
mt_data_raw <- read_mt("example.mt")

# Use read_bulk to read all raw data files ending with ".mt" that are
# stored in the folder "raw_data" (in the current working directory)
library(readbulk)
mt_data_raw <- read_bulk("raw_data", fun=read_mt, extension=".mt")

# Import the data into mousetrap
mt_data <- mt_import_wide(mt_data_raw)

## End(Not run)
```
scale_within

Scale and center variables within the levels of another variable.

Description

scale_within centers and/or scales variables in a data.frame (using scale) depending on the levels of one or more other variables. By default, variables are standardized (i.e., centered and scaled). A typical application is the within-subject standardization of variables in a repeated measures design.

Usage

scale_within(
  data,
  variables = NULL,
  within = NULL,
  prefix = "",
  center = TRUE,
  scale = TRUE
)

Arguments

data a data.frame.
variables a character string (or vector) specifying one or more variables that scale is applied to. If unspecified, scale_within will be applied to all variables in data.
within an optional character string specifying the name of one or more variables in data. If specified, scale is applied separately for each of the levels of the variable (or for each combination of levels, if more than one variable is specified). Alternatively, a vector directly containing the level values.
prefix a character string that is inserted before each scaled variable. By default (empty string) the original variables are replaced.
center argument passed on to scale.
scale argument passed on to scale.

Value

The original data.frame including the centered and / or scaled variables.

Author(s)

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See Also

circle for the R base scale function.
mt_standardize for standardizing measures in a mousetrap data object.
Examples

```r
ChickWeight_scaled <- scale_within(
  ChickWeight, variables="weight",
  within="Chick", prefix="z_")
```
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