# Package 'eyetools'

July 22, 2025

Type Package

Title Analyse Eye Data

Version 0.9.2

**Description** Enables the automation of actions across the pipeline, including initial steps of transforming binocular data and gap repair to event-based processing such as fixations, saccades, and entry/duration in Areas of Interest (AOIs). It also offers visualisation of eye movement and AOI entries. These tools take relatively raw (trial, time, x, and y form) data and can be used to return fixations, saccades, and AOI entries and time spent in AOIs. As the tools rely on this basic data format, the functions can work with data from any eye tracking device. Implements fixation and saccade detection using methods proposed by Salvucci and Goldberg (2000) <doi:10.1145/355017.355028>.

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URL https://tombeesley.github.io/eyetools/

BugReports https://github.com/tombeesley/eyetools/issues

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AOI\_seq

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### Description

Analyses the sequence of entries into defined AOI regions across trials. Can only be used with fixation data with a "fix\_n" column denoting fixation events. Assumes that AOIs are non-overlapping and hasn't been tested with overlapping AOIs. Consecutive fixations within an AOI are grouped together as a single entry. Non-consecutive fixations in the same AOI (i.e., with an intervening fixation in no AOI) are treated as two separate entries.

```
AOI_seq(data, AOIs, AOI_names = NULL, progress = TRUE)
```

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#### **Arguments**

data	A dataframe with fixation data (from fixation_dispersion). Either single or multi participant data
AOIs	A dataframe of areas of interest (AOIs), with one row per AOI $(x, y, width\_radius, height)$ .
AOI_names	An optional vector of AOI names to replace the default "AOI_1", "AOI_2", etc.
progress	Display a progress bar

### Value

a dataframe containing the sequence of entries into AOIs on each trial, entry/exit/duration time into AOI

### **Examples**

```
data <- combine_eyes(HCL)
fix_d <- fixation_dispersion(data)

AOI_seq(fix_d, AOIs = HCL_AOIs)</pre>
```

AOI\_time

Analysis of time spent in areas of interest

### Description

Analyses total time on defined AOI regions across trials. Works with fixation and raw data as the input (must use one or the other, not both).

```
AOI_time(
  data,
  data_type = NULL,
  AOIs,
  AOI_names = NULL,
  sample_rate = NULL,
  as_prop = FALSE,
  trial_time = NULL
)
```

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#### **Arguments**

data	A dataframe of either fixation data (from fix_dispersion) or raw data
data_type	Whether data is a fixation ("fix") or raw data ("raw")
AOIs	A dataframe of areas of interest (AOIs), with one row per AOI (x, y, width_radius, height).
AOI_names	An optional vector of AOI names to replace the default "AOI_1", "AOI_2", etc.
sample_rate	Optional sample rate of the eye-tracker (Hz) for use with data. If not supplied, the sample rate will be estimated from the time column and the number of samples.
as_prop	whether to return time in AOI as a proportion of the total time of trial
trial_time	needed if as_prop is set to TRUE. a vector of the time taken in each trial. Equal to the length of x trials by y participants in the dataset

#### **Details**

Analyses data separately for each unique combination of values in pID and trial. Returned values can be absolute time or proportion of time over the period.

#### Value

a dataframe containing the time on the passed AOIs for each trial. One column for each AOI separated by trial.

AOI\_time\_binned 5

AOI_time_binned Binned time analysis of area of interest entries	
--	--

#### **Description**

Analyses total time on defined AOI regions across trials separated into bins. Works with raw data as the input. Data can be separated into bins of a given length of time and the number of bins per trial is calculated automatically, keeping the bin length consistent across varying lengths of trial. Any data that cannot fill a bin (typically the last few milliseconds of the trial) are dropped to ensure that bins are of a consistent length

### Usage

```
AOI_time_binned(
    data,
    AOIs,
    AOI_names = NULL,
    sample_rate = NULL,
    bin_length = NULL,
    max_time = NULL,
    as_prop = FALSE
)
```

#### **Arguments**

data	A dataframe of raw data
AOIs	A dataframe of areas of interest (AOIs), with one row per AOI $(x, y, width\_radius, height)$ .
AOI_names	An optional vector of AOI names to replace the default "AOI_1", "AOI_2", etc.
sample_rate	Optional sample rate of the eye-tracker (Hz) for use with data. If not supplied, the sample rate will be estimated from the time column and the number of samples.
bin_length	the time duration to be used for each bin.
max_time	maximum length of time to use, default is total trial length
as_prop	whether to return time in AOI as a proportion of the total time of trial

#### **Details**

AOI\_time\_binned can take either single participant data or multiple participants, where participants are demarcated by values in the "pID" column.

#### Value

a dataframe containing the time on the passed AOIs for each trial. One column for each AOI separated by trial.

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#### **Examples**

```
data <- combine_eyes(HCL)

#with bins of 100ms each and only for the first 2000ms
AOI_time_binned(data = data, AOIs = HCL_AOIs,
    bin_length = 100, max_time = 2000)</pre>
```

combine\_eyes

Combine binocular data into single X/Y coordinate pairs

#### **Description**

Combines the data from binocular samples into X/Y coordinate pairs. Two methods can be used: "average" or "best\_eye". For "average", the result is based on the average of the two eyes for each sample, or for samples where there is data from only a single eye, that eye is used. For "best\_eye", a summary of the proportion of missing samples is computed, and the eye with the fewest missing samples is used.

### Usage

```
combine_eyes(data, method = "average")
```

#### **Arguments**

data raw data with columns time, left\_x, left\_y, right\_x, right\_y, and trial method either "average" or "best\_eye" - see description.

#### Value

a dataframe of x-2 variables (with left\_x and right\_x condensed to x, and left\_y and right\_y condensed to y) and the same number of observations as the input data

```
combine_eyes(HCL, method = "average")
```

compare\_algorithms 7

compare_algorithms	A battery of metrics and plots to compare the two algorithms (disper-
	sion and VTI)

#### **Description**

A tool for comparing the two different algorithms present in this package. This function is useful for assessing the data as well as exploring which algorithm is likely to fit data more appropriately. The raw data is run through both algorithms (using the same specified dispersion tolerances, etc.) before making comparisons of the underlying data. Can only be used for single participant data.

### Usage

```
compare_algorithms(
  data,
  plot_fixations = TRUE,
  print_summary = TRUE,
  sample_rate = NULL,
  threshold = 100,
  min_dur = 150,
  min_dur_sac = 20,
  disp_tol = 100,
  NA_tol = 0.25,
  smooth = FALSE
)
```

#### **Arguments**

data	A dataframe with raw data (time, x, y, trial) for one participant
${\tt plot\_fixations}$	Whether to plot the detected fixations. default as TRUE
<pre>print_summary</pre>	Whether to print the summary table. default as TRUE
sample_rate	sample rate of the eye-tracker. If default of NULL, then it will be computed from the timestamp data and the number of samples. Supplied to the VTI algorithm
threshold	velocity threshold (degrees of VA $\prime$ sec) to be used for identifying saccades. Supplied to the VTI algorithm
min_dur	Minimum duration (in milliseconds) of period over which fixations are assessed. Supplied to both algorithms.
min_dur_sac	Minimum duration (in milliseconds) for saccades to be determined. Supplied to the VTI algorithm
disp_tol	Maximum tolerance (in pixels) for the dispersion of values allowed over fixation period. Supplied to both algorithms
NA_tol	the proportion of NAs tolerated within any window of samples that is evaluated as a fixation. Supplied to the dispersion algorithm
smooth	include a call to eye tools::smoother on each trial. Supplied to the VTI algorithm

conditional\_transform

#### Value

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a list of the fixation data, correlation output, and data used for plotting

### **Examples**

```
data <- combine_eyes(HCL)
data <- interpolate(data)
compare_algorithms(data[data$pID == 119,])</pre>
```

 $conditional\_transform$   $conditional\_transform$ 

### **Description**

A function to perform conditional transformations of the x/y raw data. The function takes the dataframe and performs a single axis flip based on the values specified in the cond\_column. The primary use of this function is to correct or normalise the data when counterbalancing stimulus placement within experiments (e.g., having a target stimulus appear on the left and right equally often)

#### Usage

```
conditional_transform(
  data,
  flip = c("x", "y"),
  cond_column,
  cond_values,
  resolution_x = 1920,
  resolution_y = 1080,
  message = TRUE
)
```

#### Arguments

data	a dataframe that includes columns x and y and the column specified in cond_column. Can be raw, fixation, or saccade data.
flip	either "x", to flip across vertical midline, or "y" to flip across horizontal midline
cond_column	a column name, on which the flips are conditional
cond_values	a single value or vector stating which values in con_column result in a flip
resolution_x	screen size in pixels for the x axis
resolution_y	screen size in pixels for the y axis
message	whether to output messages during function. Useful to turn off when using in a vectorised fashion where it is running multiple times

create\_AOI\_df

#### Value

a dataframe of the equivalent format as the input data

### Examples

create\_AOI\_df

Create a blank data frame for populating with AOIs

### **Description**

Create a blank data frame for populating with AOIs

#### Usage

```
create_AOI_df(num_AOIs = 3, shape = "rect", AOI_data = NULL)
```

#### Arguments

num\_AOIs number of AOIs, setting the number of rows

shape whether the AOI is rectangular ("rect") or circular ("circ")

AOI\_data a list of data for each AOI, ordered by x, y, width\_radius, and height

#### Value

a dataframe in the standard format required for eyetools

10 dist\_to\_visual\_angle

### **Description**

Takes a single value or vector of distances and returns the visual angle equivalent.

#### Usage

```
dist_to_visual_angle(
  vector,
  dist_type = "cm",
  view_dist_cm = 60,
  screen_width_cm = 51,
  screen_width_pixels = 1920
)
```

#### Arguments

```
vector vector of distances (or single distance)

dist_type default is "cm". Specify "pixel" for conversion from pixel values.

view_dist_cm viewing distance in cm. Default of 60cm.

screen_width_cm used in conversion of pixel values. Default is 51 cm (24" monitor).

screen_width_pixels used in conversion of pixel values. Default is 1920 pixels.
```

#### Value

an equivalent-sized object to the input

```
# calculate visual angle for stimulus of 5cm
dist_to_visual_angle(5)

# calculate visual angle of stimuli 2 and 10cm width at 50 cm viewing angle
dist_to_visual_angle(c(2,10), view_dist_cm = 50)

# calculate visual angle of 150 pixel wide
dist_to_visual_angle(150, dist_type = "pixels")
```

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fixation\_dispersion Fixat

Fixation detection using a dispersion method

### Description

Detects fixations by assessing dispersion of the eye position, using a method that is similar to that proposed by Salvucci & Goldberg (2000). Evaluates the maximum dispersion (distance) between x/y coordinates across a window of data. Looks for sufficient periods in which this maximum dispersion is below the specified dispersion tolerance. NAs are considered breaks in the data and are not permitted within a valid fixation period.

### Usage

```
fixation_dispersion(
  data,
  min_dur = 150,
  disp_tol = 100,
  NA_tol = 0.25,
  progress = TRUE
)
```

### Arguments

data	A dataframe with raw data (time, x, y, trial) for one participant (the standardised raw data form for eyetools)
min_dur	Minimum duration (in milliseconds) of period over which fixations are assessed
disp_tol	Maximum tolerance (in pixels) for the dispersion of values allowed over fixation period
NA_tol	the proportion of NAs tolerated within any window of samples that is evaluated as a fixation
progress	Display a progress bar

#### **Details**

It can take either single participant data or multiple participants, where participants are demarcated by values in the "pID" column.

### Value

a dataframe containing each detected fixation by trial, with mean x/y position in pixel, start and end times, and duration.

### References

Salvucci, D. D., & Goldberg, J. H. (2000). Identifying fixations and saccades in eye-tracking protocols. Proceedings of the Symposium on Eye Tracking Research & Applications - ETRA '00, 71–78.

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#### **Examples**

```
data <- combine_eyes(HCL)
fixation_dispersion(data)</pre>
```

fixation\_VTI

Fixation detection using a velocity threshold identification method

#### **Description**

Determine fixations by assessing the velocity of eye-movements, using a method that is similar to that proposed by Salvucci & Goldberg (2000). Applies the algorithm used in VTI\_saccade and removes the identified saccades before assessing whether separated fixations are outside of the dispersion tolerance. If they are outside of this tolerance, the fixation is treated as a new fixation regardless of the length of saccade separating them. Compared to fixation\_dispersion(), fixation\_VTI() is more conservative in determining a fixation as smaller saccades are discounted and the resulting data is treated as a continued fixation (assuming it is within the pixel tolerance set by disp\_tol). Returns a summary of the fixations found per trial, including start and end coordinates, timing, duration, mean velocity, and peak velocity.

#### Usage

```
fixation_VTI(
  data,
  sample_rate = NULL,
  threshold = 100,
  min_dur = 150,
  min_dur_sac = 20,
  disp_tol = 100,
  smooth = FALSE,
  progress = TRUE
)
```

#### **Arguments**

data	A dataframe with raw data (time, x, y, trial) for one participant
sample_rate	sample rate of the eye-tracker. If default of NULL, then it will be computed from the timestamp data and the number of samples
threshold	velocity threshold (degrees of VA / sec) to be used for identifying saccades.
min_dur	Minimum duration (in milliseconds) of period over which fixations are assessed
min_dur_sac	Minimum duration (in milliseconds) for saccades to be determined
disp_tol	Maximum tolerance (in pixels) for the dispersion of values allowed over fixation period
smooth	include a call to eyetools::smoother on each trial
progress	Display a progress bar

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#### **Details**

Analyses data separately for each unique combination of values in pID and trial.

#### Value

a dataframe containing each detected fixation by trial, with mean x/y position in pixel, start and end times, and duration.

#### References

Salvucci, D. D., & Goldberg, J. H. (2000). Identifying fixations and saccades in eye-tracking protocols. Proceedings of the Symposium on Eye Tracking Research & Applications - ETRA '00, 71–78.

#### **Examples**

```
data <- combine_eyes(HCL)
data <- interpolate(data)
fixation_VTI(data)</pre>
```

**HCL** 

Example dataset from that contains binocular eye data from two participants from a simple contingency learning task (the data are from Beesley, Nguyen, Pearson, & Le Pelley, 2015). In this task there are two stimuli that appear simultaneously on each trial (to the left and right of the screen). Participants look at these cues and then make a decision by selecting an "outcome response" button.

#### **Description**

The dataset contains data from two participants and the first six trials of the study.

#### Usage

HCL

#### **Format**

A dataframe of 31,041 observations and seven variables

pID participant ID

time timestamp of the sample (milliseconds)

**left\_x** x coordinate of the left eye

**left\_y** y coordinate of the left eye

right\_x x coordinate of the right eye

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right\_y y coordinate of the right eye
trial trial number ...

HCL\_AOIs

Example AOIs for use with HCL

#### **Description**

This dataframe contains three rectangular areas of interest (AOIs), set out for use with the HCL dataset. Values are in pixels.

#### Usage

HCL\_AOIs

#### **Format**

A data frame with 3 rows and 4 variables:

- x centred x coordinate of the AOI
- y centred y coordinate of the AOI

width\_radius either the width of the AOI, or the radius for circular AOIs

height the height of the AOI; should be NA for circular AOIs ...

HCL\_behavioural

Example dataset of behavioural data to complement dataset HCL.

### Description

This contains information on stimuli (such as the side the predictive cue was presented on) as well as response data, including accuracy and response times

#### Usage

HCL\_behavioural

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#### **Format**

```
A dataframe of 12 observations and eight variables
```

```
pID participant ID

trial trial number

P_cue Are these necessary columns?

NP_cue Are these necessary columns?

cue_order whether the predictive cue os presented on the left (1) or the right (2)

correct_out NAre these necessary columns?

accuracy response accuracy

RT response time in milliseconds ...
```

hdf5\_get\_event

Get messgaes stored in TOBII-generated HDF5 files

### Description

A function to get the message event files from a TOBII-generated hdf5 file to dataframe. Used when a Psychopy experiment uses the io.sendMessageEvent() to record events

### Usage

```
hdf5_get_event(filename)
```

### Arguments

filename

the hdf5 file generated from TOBII

#### Value

A dataframe of message events as recorded by TOBII eye trackers

```
## Not run:
raw_data <- hdf5_get_event("example_TOBII.hdf5")
## End(Not run)</pre>
```

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hdf5\_to\_df

Convert TOBII-generated HDF5 files to dataframe

### **Description**

A function to convert TOBII-generated hdf5 files to a dataframe

#### Usage

```
hdf5_to_df(filename)
```

### Arguments

filename

the hdf5 file generated from TOBII

#### Value

A list of dataframes collected from the eyetracker content, if only one eyetracking event is present, return this as a single dataframe

#### **Examples**

```
## Not run:
raw_data <- hdf5_to_df("example_TOBII.hdf5")
## End(Not run)</pre>
```

interpolate

Interpolation of missing data (NAs)

#### **Description**

Extends the zoo::na.approx and zoo::na.spline functions to include a report which provides the proportion of missing data before and after the interpolation process. This is handy for evaluating the effectiveness of the repair.

```
interpolate(
  data,
  maxgap = 150,
  method = "approx",
  sample_rate = NULL,
  report = FALSE
)
```

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### **Arguments**

data	dataframe with columns time, $x$ , $y$ , trial (the standardised raw data form for eyeproc)
maxgap	maximum time gap of consecutive trackloss to fill (in ms). Any longer gaps will be left unchanged (see zoo package)
method	"approx" for linear interpolation or "spline" for cubic spline interpolation
sample_rate	Optional sample rate of the eye-tracker (Hz) for use with data. If not supplied, the sample rate will be estimated from the time column and the number of samples.
report	default is FALSE. If TRUE, then the return value is a list containing the returned

Analyses data separately for each unique combination of values in pID and trial.

data frame and the report.

#### Value

**Details** 

a dataframe of the same shape of the input data

### **Examples**

```
data <- combine_eyes(HCL)
interpolate(data, maxgap = 150)</pre>
```

plot\_AOI\_growth

Plots absolute or proportional time spent in AOIs over time

## Description

A visualisation tool for plotting the changes in defined AOI regions across a single trial time.

```
plot_AOI_growth(
  data = NULL,
  pID_values = NULL,
  trial_values = NULL,
  AOIs = NULL,
  AOI_names = NULL,
  type = "abs",
  plot_time_not_in_AOI = FALSE
)
```

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### **Arguments**

raw data in standard raw data form (time, x, y, trial)

pID\_values specify particular values within 'pID' to plot data from certain participants

trial\_values can be used to select particular trials within the data

AOIs A dataframe of areas of interest (AOIs), with one row per AOI (x, y, width\_radius, height)

AOI\_names An optional vector of AOI names to replace the default "AOI\_1", "AOI\_2", etc.

To omit AOIs from the plot, use NA in relevant vector position

type either "abs" (absolute) or "prop" (proportion)

plot\_time\_not\_in\_AOI

boolean as to whether to include proportion of time spent outside AOIs

#### Value

a plot of the raw data

#### **Examples**

```
data <- combine_eyes(HCL)
data <- interpolate(data)
# plot absolute and then proportional
plot_AOI_growth(data = data, AOIs = HCL_AOIs, type = "abs")
plot_AOI_growth(data = data, AOIs = HCL_AOIs, type = "prop")</pre>
```

plot\_heatmap

Plot heatmap of raw data

### **Description**

Plots a heatmap of raw data.

```
plot_heatmap(
  data = NULL,
  pID_values = NULL,
  trial_values = NULL,
  bg_image = NULL,
  res = c(0, 1920, 0, 1080),
  flip_y = FALSE,
  plot_type = "density",
  alpha_range = c(0.1, 0.8),
  plot_header = FALSE
)
```

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#### Arguments

data data in standard raw data form (time, x, y, trial) specify particular values within 'pID' to plot data from certain participants pID\_values trial values specify particular values within 'trial' to plot data from certain trials bg\_image The filepath of a PNG image to be added to the plot, for example to show a screenshot of the task. resolution of the display to be shown, as a vector (xmin, xmax, ymin, ymax) res reverse the y axis coordinates (useful if origin is top of the screen) flip\_y Specify the nature of the data displayed. Either "density" (default) or "hex" plot\_type a pair of values between 0 and 1. The first is a cut off, whereby lower values are alpha\_range not displayed. The second value sets the transparancy of the visible poitns. display the header title text which explains graphical features of the plot. plot\_header

#### Value

a plot of the raw data

#### **Examples**

```
data <- combine_eyes(HCL)
# plot all trials data
plot_heatmap(data, pID_values = 118, alpha_range = c(0.3,0.8))
#plot one trial
plot_heatmap(data, trial_values = 1)</pre>
```

plot\_seq

Plot of raw data over time

#### **Description**

A tool for visualising the timecourse of raw data over a single trial. Participant and trial values can be selected from the data. If values for these parameters are not provided then a single participant and a single trial will be sampled at random. Data can be split into bins by time or by the number of bins.

```
plot_seq(
  data = NULL,
  AOIs = NULL,
  trial_values = NULL,
  pID_values = NULL,
```

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```
bg_image = NULL,
res = c(0, 1920, 0, 1080),
flip_y = FALSE,
plot_header = FALSE,
bin_time = NULL,
bin_range = NULL
)
```

### Arguments

data	A dataframe with raw data. If multiple trials are used, then one trial is sampled at random.
AOIs	A dataframe of areas of interest (AOIs), with one row per AOI $(x, y, width\_radius, height)$ .
trial_values	specify particular values within 'trial' to plot data from certain trials
pID_values	specify particular values within 'pID' to plot data from certain participants
bg_image	The filepath of a PNG image to be added to the plot, for example to show a screenshot of the task.
res	resolution of the display to be shown, as a vector (xmin, xmax, ymin, ymax)
flip_y	reverse the y axis coordinates (useful if origin is top of the screen)
plot_header	display the header title text which explains graphical features of the plot.
bin_time	if wanting to split data into bins, the time (in ms) for each bin of data to be displayed
bin_range	if wanting to split data into bins, the first and last bin to be display, e.g., c(1,5)

### Value

a plot of the raw data representing changes over time

```
data <- combine_eyes(HCL)

# plot the raw data
plot_seq(data)

# with AOIs
plot_seq(data, AOIs = HCL_AOIs)

# plot raw data with bins
plot_seq(data, bin_time = 500)</pre>
```

plot\_spatial 21

plot_spatial Plot raw data and fixations
--

### Description

A tool for visualising raw eye-data, processed fixations, and saccades. Can use all three data types together and independently. Fixations can be labeled in the order they were made. Can overlay areas of interest (AOIs) and customise the resolution.

### Usage

```
plot_spatial(
  raw_data = NULL,
  fix_data = NULL,
  sac_data = NULL,
  AOIs = NULL,
  pID_values = NULL,
  trial_values = NULL,
  bg_image = NULL,
  res = c(0, 1920, 0, 1080),
  flip_y = FALSE,
  show_fix_order = TRUE,
  plot_header = FALSE
)
```

### Arguments

raw_data	data in standard raw data form (time, x, y, trial)
fix_data	data output from fixation function
sac_data	data output from saccade function
AOIs	A dataframe of areas of interest (AOIs), with one row per AOI (x, y, width_radius, height). If using circular AOIs, then the 3rd column is used for the radius and the height should be set to NA.
pID_values	specify particular values within 'pID' to plot data from certain participants
trial_values	specify particular values within 'trial' to plot data from certain trials
bg_image	The filepath of a PNG image to be added to the plot, for example to show a screenshot of the task.
res	resolution of the display to be shown, as a vector (xmin, xmax, ymin, ymax)
flip_y	reverse the y axis coordinates (useful if origin is top of the screen)
show_fix_order	label the fixations in the order they were made
plot_header	display the header title text which explains graphical features of the plot.

### Value

```
a plot of the raw data
```

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#### **Examples**

```
data <- combine_eyes(HCL)
# plot the raw data
plot_spatial(raw_data = data, pID_values = 118)

# plot both raw and fixation data together
plot_spatial(raw_data = data, fix_data = fixation_dispersion(data), pID_values = 118)

#plot one trial
plot_spatial(raw_data = data, fix_data = fixation_dispersion(data), trial_values = 6)</pre>
```

saccade\_VTI

Velocity threshold identification of saccades

#### **Description**

Use the velocity threshold algorithm from Salvucci & Goldberg (2000) to determine saccadic eye movements. Returns a summary of the saccades found per trial, including start and end coordinates, timing, duration, mean velocity, and peak velocity.

#### Usage

```
saccade_VTI(data, sample_rate = NULL, threshold = 150, min_dur = 20)
```

#### **Arguments**

A dataframe with raw data (time, x, y, trial) for one participant

sample\_rate sample rate of the eye-tracker. If default of NULL, then it will be computed from the timestamp data and the number of samples

threshold velocity threshold (degrees of VA / sec) to be used for identifying saccades

min\_dur minimum duration (ms) expected for saccades. This helps to avoid identification of very short saccades occurring at the boundary of velocity threshold

### Details

Analyses data separately for each unique combination of values in pID and trial.

#### Value

a data frame giving the saccades found by trial

#### References

Salvucci, D. D., & Goldberg, J. H. (2000). Identifying fixations and saccades in eye-tracking protocols. Proceedings of the Symposium on Eye Tracking Research & Applications - ETRA '00, 71–78.

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### **Examples**

```
data <- combine_eyes(HCL)
saccade_VTI(data)</pre>
```

smoother

Smoothing of raw data

### Description

A wrapper for the stats::loess function, with default parameters suitable for smoothing raw eye data

### Usage

```
smoother(data, span = 0.05, plot = FALSE)
```

### Arguments

data	A dataframe with raw data (time, x, y, trial) for one participant
span	From stats::loess. The parameter alpha which controls the degree of smoothing.
plot	whether to plot the raw and smoothed plot for inspection

#### **Details**

Analyses data separately for each unique combination of values in pID and trial.

### Value

a dataframe of the same shape as the input data

```
data <- combine_eyes(HCL)
smoother(data)

#with an inspection plot
smoother(data, span = .02, plot = TRUE)</pre>
```

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