

Package ‘webmorphR’

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Title Reproducible Stimuli

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Description Create reproducible image stimuli,
specialised for face images with 'psychomorph' or 'webmorph' templates.

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URL <https://debruine.github.io/webmorphR/>,
<https://github.com/debruine/webmorphR>

BugReports <https://github.com/debruine/webmorphR/issues>

Depends R (>= 4.1.0)

Imports tools, grDevices, stats, dplyr, jsonlite, magick, geomorph,
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<code>add_info</code>	<i>Add Information</i>
-----------------------	------------------------

Description

Add info with a data table that contains the info in either the same order as the stimulus list, or matching the stimuli item name with the column specified by `.by`.

Usage

```
add_info(stimuli, ..., .by = NULL)
```

Arguments

<code>stimuli</code>	list of stimuli
<code>...</code>	data table or named vectors of info to add
<code>.by</code>	the column to use to match info to stimuli names; leave NULL if the data are to be matched by order

Details

You can also add data as named vectors.

Value

list of stimuli with info added

See Also

Other info: [compare\(\)](#), [get_info\(\)](#), [get_point\(\)](#), [height\(\)](#), [metrics\(\)](#), [rename_stim\(\)](#), [width\(\)](#)

Examples

```
stimuli <- demo_stim() |>
  add_info(project = "XXX", gender = c("F", "M"))

stimuli$f_multi$info |> str()
```

align

Align templates and images

Description

Align images so that template points line up. Defaults to two-point alignment of the first two points in your template (usually the eyes) to their mean coordinate position across the stimuli.

Usage

```
align(
  stimuli,
  pt1 = 0,
  pt2 = 1,
  x1 = NULL,
  y1 = NULL,
  x2 = NULL,
  y2 = NULL,
  width = NULL,
  height = NULL,
  ref_img = NULL,
  fill = wm_opts("fill"),
  procrustes = FALSE
)
```

Arguments

stimuli	list of stimuli
pt1	The first point to align (defaults to 0)
pt2	The second point to align (defaults to 1)
x1, y1, x2, y2	The coordinates to align the first and second point to
width, height	The dimensions of the aligned images
ref_img	The reference image to get coordinates and dimensions from if they are NULL
fill	background color if cropping goes outside the original image, see color_conv()
procrustes	logical; whether to use procrustes alignment

Details

Setting pt1 the same as pt2 aligns 1 point, but does not resize or rotate images. Setting pt1 and pt2 aligns 2 points, resizing and rotating faces. Setting procrustes = TRUE uses Procrustes analysis to resize and rotate images to be as close as possible to a mean shape.

You can specify the x and y coordinates to align, and the width and height of the output images, or set them from a reference image. The reference image (ref_img) can be a stim, a 1-item stimlist, or the index or name of a stim in stimuli. It defaults to average of all stimuli if NULL.

Visualise the template points with `draw_tem()` to determine which to align, using `pt.shape = "index"`.

Value

list of stimuli with aligned images and/or templates

See Also

Stimulus manipulation functions `crop_tem()`, `crop()`, `greyscale()`, `horiz_eyes()`, `image_func()`, `mask_oval()`, `mask()`, `mirror()`, `pad()`, `resize()`, `rotate()`, `to_size()`

Examples

```
# align eye points to specific x and y coordinates
# in a 300x300 pixel image
demo_unstandard(1:3) |>
  align(pt1 = 0, pt2 = 1,
        x1 = 100, x2 = 200, y1 = 100, y2 = 100,
        width = 300, height = 300)

orig <- demo_unstandard()

# align to bottom-centre of nose (average position)
align(orig, pt1 = 55, pt2 = 55, fill = "dodgerblue")

# align to pupils of second image
align(orig, ref_img = 2, fill = "dodgerblue")

## Not run:
# procrustes align to average position
# this requires XQuartz on mac and may not run on linux
align(orig, procrustes = TRUE, fill = "dodgerblue")

## End(Not run)
```

animate *Create an animated gif from a list of stimuli*

Description

Create an animated gif from a list of stimuli

Usage

```
animate(stimuli, fps = 1, loop = 0, rev = FALSE)
```

Arguments

stimuli	list of stimuli
fps	frames per second
loop	how many times to loop the animation (0 = infinite)
rev	whether to loop back and forth (TRUE) or in one direction (FALSE)

Value

magick image

See Also

Stimulus creating functions [as_stimlist\(\)](#), [blank\(\)](#), [new_stimlist\(\)](#), [new_stim\(\)](#), [read_img\(\)](#), [read_stim\(\)](#), [read_tem\(\)](#), [write_stim\(\)](#)

Examples

```
# slideshow of images (1/second)
demo_stim() |> animate()

# rotate a face
degrees <- seq(0, 350, 10)
demo_stim(1) |>
  mask() |>
  rep(length(degrees)) |>
  rotate(degrees) |>
  animate(fps = 10)
```

`as_ggplot`*Convert stimuli to a ggplot*

Description

Convert a stimulus or list of stimuli into a ggplot, which can be further used with ggplot functions.

Usage

```
as_ggplot(stimuli, ...)
```

Arguments

<code>stimuli</code>	list of stimuli
<code>...</code>	Additional arguments to pass to <code>plot_stim()</code> if stimuli contains more than 1 image

Value

a ggplot object

See Also

Visualisation functions `draw_tem()`, `gglabel()`, `label()`, `mlabel()`, `plot.stimlist()`, `plot.stim()`, `plot_rows()`, `plot_stim()`

Examples

```
stimuli <- demo_stim()
gg <- as_ggplot(stimuli)

# add to ggplot object; coordinates are pixels
# (images are 500x500 each, plus 10px padding)
gg +
  ggplot2::geom_vline(xintercept = 0, color = "red") +
  ggplot2::geom_vline(xintercept = 1030, color = "blue") +
  ggplot2::geom_hline(yintercept = 0, color = "green") +
  ggplot2::geom_hline(yintercept = 520, color = "purple") +
  ggplot2::annotate("point", x = 515, y = 260, size = 10) +
  ggplot2::labs(
    title = "This is a ggplot!",
    caption = "Made with webmorphR"
  )
```

`as_stimlist`*Convert list to stimlist*

Description

Checks if an object is a stimulus or list of stimuli and repairs common problems.

Usage

```
as_stimlist(x)
```

Arguments

`x` The object

Details

Some webmorphR functions, like `plot()` and `print()` require objects to have a "stimlist" class. If you've processed a list of stimuli with iterator functions like `lapply()` or `purrr::map()` and the resulting object prints or plots oddly, it is probably unclassed, and this function will fix that.

Value

A stimlist

See Also

Stimulus creating functions `animate()`, `blank()`, `new_stimlist()`, `new_stim()`, `read_img()`, `read_stim()`, `read_tem()`, `write_stim()`

Examples

```
stimuli <- demo_stim() |>
  lapply(function(stim) {
    # remove template lines
    stim$lines <- NULL
    return(stim)
  })

class(stimuli)

## Not run:
plot(stimuli) # error

## End(Not run)

s <- as_stimlist(stimuli)
class(s)
plot(s)
```

auto_delin	<i>Auto-Delineation</i>
------------	-------------------------

Description

Automatically delineate faces using Face++ (an external service). Since each delineation counts against a daily limit, you need to set up your own Face++ account (see details below).

Usage

```
auto_delin(stimuli, model = c("fpp106", "fpp83"), replace = FALSE, face = 1)
```

Arguments

stimuli	list of stimuli
model	Which model (fpp106, fpp83)
replace	logical; whether to replace original templates - if FALSE, only gets templates for images with no template
face	which face to delineate in each image if there is more than 1

Details

To use Face++ auto-delineation, you need to get your own free API key from <https://www.faceplusplus.com>. After signing up for an account, go to <https://console.faceplusplus.com/app/apikey/list> and request a free API key. Add the key and secret to your .Renviron file as follows:

```
FACEPLUSPLUS_KEY="1234567890abcdefghijk"  
FACEPLUSPLUS_SECRET="1234567890abcdefghijk"
```

Value

list of stimuli with templates

See Also

Template functions [average_tem\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Examples

```
## Not run:  
# requires an API key in .Renviron  
auto_fpp106 <- demo_stim() |>  
  auto_delin(model = "fpp106", replace = TRUE)  
  
## End(Not run)
```

average_tem	<i>Average templates</i>
-------------	--------------------------

Description

This function just averages the templates. An average image is returned, but it is just all the images superimposed. To create a template-aware average, see [avg\(\)](#).

Usage

```
average_tem(stimuli, name = "average")
```

Arguments

stimuli	list of stimuli
name	Name for the average

Value

list of stimuli consisting of just the average

See Also

Template functions [auto_delin\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Examples

```
tem_only_avg <- demo_stim() |> average_tem()

# view the average template
draw_tem(tem_only_avg, bg = "white")

# view the superimposed image
tem_only_avg
```

avg	<i>Average Images</i>
-----	-----------------------

Description

Create an average from a list of delineated stimuli.

Usage

```
avg(
  stimuli,
  texture = TRUE,
  norm = c("none", "twopoint", "rigid"),
  normpoint = 0:1
)
```

Arguments

stimuli	list of stimuli to average
texture	logical; whether textured should be averaged
norm	how to normalise; see Details
normpoint	points for twopoint normalisation

Details**Normalisation options:**

- none: averages will have all coordinates as the mathematical average of the coordinates in the component templates
- twopoint: all images are first aligned to the 2 alignment points designated in normpoint. Their position is set to their position in the first image in stimuli
- rigid: procrustes aligns all images to the position of the first image in stimuli

Texture:

This applies a representative texture to the average, resulting in composite images with more realistic texture instead of the very smooth, bland texture most other averaging programs create. See the papers below for methodological details.

B. Tiddeman, M. Stirrat and D. Perrett (2005). Towards realism in facial prototyping: results of a wavelet MRF method. *Theory and Practice of Computer Graphics*.

B. Tiddeman, D.M. Burt and D. Perrett (2001). *Computer Graphics in Facial Perception Research*. *IEEE Computer Graphics and Applications*, 21(5), 42-50.

Value

list of stimuli with the average image and template

See Also

WebMorph.org functions [continuum\(\)](#), [loop\(\)](#), [symmetrize\(\)](#), [trans\(\)](#), [webmorph_up\(\)](#)

Examples

```
if (webmorph_up()) {
  demo_stim() |> avg()
}
```

blank	<i>Make blank images</i>
-------	--------------------------

Description

Make blank images

Usage

```
blank(n = 1, width = 100, height = 100, color = "white", names = "img")
```

Arguments

n	the number of images to return
width, height	image dimensions
color	background color, see color_conv()
names	names of the images (appended with index if < n)

Value

list of stimuli

See Also

Stimulus creating functions [animate\(\)](#), [as_stimlist\(\)](#), [new_stimlist\(\)](#), [new_stim\(\)](#), [read_img\(\)](#), [read_stim\(\)](#), [read_tem\(\)](#), [write_stim\(\)](#)

Examples

```
stimuli <- blank(5, 100, 250, color = rainbow(5))  
label(stimuli, size = 20)
```

bounds	<i>Get template bounds</i>
--------	----------------------------

Description

Get template bounds

Usage

```
bounds(stimuli, each = FALSE)
```

Arguments

stimuli	A stimlist
each	Whether to calculate max and min for the full set (default) or each image separately

Value

A list of min and max x and y values

Examples

```
demo_stim() |> bounds() |> str()
demo_stim() |> bounds(each = TRUE)
```

centroid	<i>Get center coordinates</i>
----------	-------------------------------

Description

Get center coordinates

Usage

```
centroid(stimuli, points = NULL)
```

Arguments

stimuli	list of stimuli
points	which points to include (0-based); if NULL, all points will be used

Value

named matrix of centroid x and y coordinates

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Examples

```
demo_stim() |> centroid()

# get the centre of the eye points
demo_stim() |> centroid(0:1)
```

change_lines	<i>Change template lines</i>
--------------	------------------------------

Description

Alter, add or remove lines in a template

Usage

```
change_lines(stimuli, line_id = 1, pts = NULL)
```

Arguments

stimuli	list of stimuli
line_id	index of the line to change
pts	vector of points to change the line_idx to (deletes line if NULL)

Value

stimlist with altered templates

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [centroid\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Examples

```
# get image with dlib70 template and view lines
s <- demo_tems("dlib70")
s[[1]]$lines

# remove all lines
s2 <- change_lines(s, line_id = 1:13, pts = NULL)
s2[[1]]$lines

# visualise point indices
draw_tem(s2, pt.shape = "index", pt.size = 15)

# add a new line
s3 <- change_lines(s2, line_id = "face_outline",
                  pts = c(2:18, 28:19, 2))
s3[[1]]$lines
draw_tem(s3)
```

`color_conv`*Convert colors*

Description

Convert from common color inputs to specified output type, adding alpha transparency for output formats that support it (hexa, rgba).

Usage

```
color_conv(  
  color,  
  alpha = 1,  
  from = c("guess", "col", "hex", "hexa", "hex3", "rgb", "rgba", "lab"),  
  to = c("hexa", "hex", "rgba", "rgb", "lab", "hsv")  
)
```

Arguments

<code>color</code>	A color in one of the input formats (see Details)
<code>alpha</code>	Alpha transparency (values ≤ 1 converted to 0-255); ignored if color has alpha already
<code>from, to</code>	Input and output color spaces, see Details below.

Details

- `color`: one of the R colours listed in `grDevices::colors()`, e.g., "red"
- `hex`: hexadecimal string, e.g., "#FF0000"
- `hexa`: hexadecimal string with alpha, e.g., "#FF0000FF"
- `hex3`: abbreviated hexadecimal string, e.g., "#F00"
- `rgb`: vector of red, green and blue values 0-255, e.g., `c(255, 0, 0)`
- `rgba`: vector of red, green, blue and alpha values 0-255, e.g., `c(255, 0, 0, 255)`
- `lab`: CIE-Lab color
- `hsv`: vector of hue, saturation and value values (0-1), e.g., `c(h=0, s = 1, v = 1)`

Value

color in to format

See Also

Other color: `col2lab()`, `lab2rgb()`

Examples

```
# different ways to input red
color_conv("red")
color_conv("#FF0000")
color_conv("#FF0000FF")
color_conv(c(255,0,0))
color_conv("rgb(255,0,0)") # you can use CSS-style text
color_conv(c(255,0,0,255))

# Lab must have names or use text format to be guessed
color_conv(c(l = 53.2, a = 80.1, b = 67.2))
color_conv("lab(53.2,80.1,67.2)")

# else, it will be guessed as rgb; fix by setting from explicitly
color_conv(c(53.2, 80.1, 67.2))
color_conv(c(53.2, 80.1, 67.2), from = "lab")

# add 50% alpha transparency to dodgerblue
color_conv("dodgerblue", alpha = 0.5, to = "rgba")
```

 compare

Image Comparison

Description

This is just a convenient way to use `magick::compareare` with `webmorph` stimuli. It defaults to the "MSE" metric, which gives a linearly increasing score to images along a morph continuum.

Usage

```
compare(stimuli, ref_stim, metric = "MSE", fuzz = 0, scale = FALSE)
```

Arguments

<code>stimuli</code>	Stimuli to compare to the <code>ref_stim</code>
<code>ref_stim</code>	A stim, 1-item stimlist, or the name or index of the comparison item in stim
<code>metric</code>	string with a metric from <code>magick::metric_types()</code> : "Undefined", "AE", "Fuzz", "MAE", "MEPP", "MSE", "NCC", "PAE", "PHASH", "PSNR", "RMSE"
<code>fuzz</code>	relative color distance (value between 0 and 100) to be considered similar in the filling algorithm (only useful for AE)
<code>scale</code>	whether to scale the values so that the maximum value is 1 and the minimum is 0 (only useful when stim is more than 1 image and includes <code>ref_stim</code>)

Details

Metric Types

- Undefined: ?
- AE: Absolute Error
- Fuzz: ?
- MAE: Mean Absolute Error
- MEPP: Mean Error Per Pixel
- MSE: Mean Squared Error
- NCC: Normalized Cross Correlation
- PAE: Peak Absolute Error
- PHASH: Perceptual Hash
- PSNR: Peak Signal-to-Noise Ratio
- RMSE: Root Mean Squared Error

How these metrics behave when comparing a morph continuum to its first image.

Increases with morph distance:

- very strong negative exponential decay at 0 fuzz; more linear with higher fuzz: AE
- strong negative exponential decay: PAE
- slight negative exponential decay: Fuzz, RMSE
- linear: MAE, MEPP, MSE
- no idea: PHASH

Decreases with morph distance:

- linear: NCC, Undefined
- slight exponential decay: PSNR

Value

Difference metric

See Also

Other info: [add_info\(\)](#), [get_info\(\)](#), [get_point\(\)](#), [height\(\)](#), [metrics\(\)](#), [rename_stim\(\)](#), [width\(\)](#)

Examples

```
stimuli <- demo_stim()
compare(stimuli, stimuli$m_multi)
compare(stimuli, stimuli$m_multi, "AE")
compare(stimuli, stimuli$m_multi, "AE", fuzz = 5)
```

continuum

Morph between two images

Description

Morph from one image to another in the specified steps.

Usage

```
continuum(from_img, to_img, from = 0, to = 1, by = 0.1, ...)
```

Arguments

from_img	image to start at
to_img	image to end at
from	starting percentage
to	ending percentage
by	step size
...	arguments to pass to trans()

Value

a list of stimuli containing each step of the continuum

See Also

WebMorph.org functions [avg\(\)](#), [loop\(\)](#), [symmetrize\(\)](#), [trans\(\)](#), [webmorph_up\(\)](#)

Examples

```
if (webmorph_up()) {  
  stimuli <- demo_stim()  
  cont <- continuum(stimuli$f_multi, stimuli$m_multi)  
  
  # create an animated gif  
  animate(cont, fps = 10, rev = TRUE)  
}
```

crop *Crop images and templates*

Description

Remove or add margins to images and templates.

Usage

```
crop(
    stimuli,
    width = 1,
    height = 1,
    x_off = NULL,
    y_off = NULL,
    fill = wm_opts("fill")
)
```

Arguments

stimuli	list of stimuli
width, height	dimensions of cropped image in pixels or proportion (<2)
x_off, y_off	offset in pixels or proportion (<1) (NULL centers cropped image)
fill	background color if cropping goes outside the original image, see color_conv()

Details

Dimensions and offsets can be set in pixels or proportions. For width and height, values less than 2 will be interpreted as proportions, otherwise pixels. For x_off and y_off, values between -1 and 1 are interpreted as proportions, otherwise pixels.

Cropping is anchored at the image center (or calculated template centroid if there is no image) unless x_off or y_off are set.

Value

list of stimuli with cropped tems and/or images

See Also

Stimulus manipulation functions [align\(\)](#), [crop_tem\(\)](#), [greyscale\(\)](#), [horiz_eyes\(\)](#), [image_func\(\)](#), [mask_oval\(\)](#), [mask\(\)](#), [mirror\(\)](#), [pad\(\)](#), [resize\(\)](#), [rotate\(\)](#), [to_size\(\)](#)

Examples

```
stimuli <- demo_stim()

# crop to 60% width and 80% height (centered)
crop(stimuli, width = .60, height = .80)

# crop to upper right quadrant
crop(stimuli, .5, .5, x_off = .5, y_off = 0)

# negative offset with fill
crop(stimuli, 260, 260,
     x_off = -10, y_off = -10,
     fill = c("red", "dodgerblue"))
```

crop_tem

Crop to template boundaries and pad

Description

Calculate the maximum and minimum x and y coordinates across the stimuli (or for each stimulus) and crop all image to this plus padding.

Usage

```
crop_tem(
  stimuli,
  top = 10,
  right = top,
  bottom = top,
  left = right,
  each = FALSE,
  ...
)
```

Arguments

stimuli	list of stimuli
top, right, bottom, left	numeric; number of pixels or proportion (<1) to pad each side
each	logical; Whether to calculate bounds for the full set (default) or each image separately
...	additional arguments to pass to <code>crop()</code>

Value

list of stimuli

See Also

Stimulus manipulation functions [align\(\)](#), [crop\(\)](#), [greyscale\(\)](#), [horiz_eyes\(\)](#), [image_func\(\)](#), [mask_oval\(\)](#), [mask\(\)](#), [mirror\(\)](#), [pad\(\)](#), [resize\(\)](#), [rotate\(\)](#), [to_size\(\)](#)

Examples

```
stimuli <- demo_stim()
ctem <- crop_tem(stimuli, each = TRUE)
draw_tem(ctem)

# demo with different templates
stimuli <- demo_tems()

# default 10 pixels around maximum template
crop_tem(stimuli)

# crop specific to each image
crop_tem(stimuli, each = TRUE)
```

delin

Manually delineate images

Description

Adjust the templates in a shiny interface. This will overwrite existing templates.

Usage

```
delin(stimuli)
```

Arguments

stimuli list of stimuli

Value

list of stimuli with new templates

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Examples

```
if (interactive()) {  
  # adjust existing delineations  
  stimuli <- demo_stim() |> delin()  
  
  # create new delineations from scratch  
  stimuli <- demo_stim() |> remove_tems() |> delin()  
}
```

demo_stim

Demo Stimuli

Description

A convenience function to get demo stimuli. See the Details below for citation and license info.

Usage

```
demo_stim(pattern = NULL)  
  
demo_tems(pattern = NULL)  
  
demo_unstandard(pattern = NULL)
```

Arguments

pattern Vector of patterns to use to search for files, or a vector of image indices (e.g., 1:4 selects the first 4 images and their templates)

Details

- `demo_stim()`: two composite faces with frl delineations; 500x500 pixels
- `demo_tems()`: an image with 5 different delineations; 675x900 pixels
- `demo_unstandard()`: a set of 10 composite faces with frl delineations; rotated, resized, and cropped so face position is not standard and each image is a different size (444 to 645 pixels)

Citation:

The images from `demo_stim()` and `demo_unstandard()` are usable on a CC-BY license, citing:
DeBruine, L. (2016). Young adult composite faces (Version 1). figshare. doi:10.6084/m9.figshare.4055130.v1

The image from `demo_tems()` is Lisa DeBruine (the author of `webmorphR`) and available on a CC-O license (no attribution needed).

Value

list of stimuli

Examples

```

demo_stim() |> label()

# visualise templates
demo_tems() |>
  draw_tem(pt.size = 10) |>
  label() |>
  plot(maxwidth = 1000)

# visualise keeping relative sizes
demo_unstandard() |>
  to_size(keep_rels = TRUE) |>
  pad(80, 0, 0, 0) |>
  label() |>
  plot(nrow = 2, maxwidth = 1000)

```

draw_tem	<i>Draw template</i>
----------	----------------------

Description

Visualise a template on an image.

Usage

```

draw_tem(
  stimuli,
  pt.color = wm_opts("pt.color"),
  pt.alpha = 0.75,
  pt.size = NULL,
  pt.shape = c("circle", "cross", "index"),
  line.color = wm_opts("line.color"),
  line.alpha = 0.5,
  line.size = NULL,
  bg = "image"
)

```

Arguments

`stimuli` list of stimuli

`pt.color`, `line.color` line or point color, see [color_conv\(\)](#)

`pt.alpha`, `line.alpha` transparency (0-1), ignored if color is a hex value with transparency. Set alpha to 0 to omit lines or points.

pt.size, line.size	size in pixels (scales to image size if NULL)
pt.shape	the shape of the points ("circle", "cross", "index")
bg	background color ("image" uses the original image)

Details

Visualising the index of each point isn't great yet and will overlay

Value

list of stimuli with template images

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [features\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Visualisation functions [as_ggplot\(\)](#), [gglabel\(\)](#), [label\(\)](#), [mlabel\(\)](#), [plot.stimlist\(\)](#), [plot.stim\(\)](#), [plot_rows\(\)](#), [plot_stim\(\)](#)

Examples

```
# get an image with 2 different templates
stimuli <- demo_tems("fr1|fpp106")
```

```
# default template
draw_tem(stimuli)
```

```
# custom template
draw_tem(stimuli,
  pt.shape = "cross",
  pt.color = "red",
  pt.alpha = 1,
  pt.size = 15,
  line.color = rgb(0, 0, 0),
  line.alpha = 0.5,
  line.size = 5)
```

```
# indexed template
draw_tem(stimuli,
  pt.shape = "index",
  pt.size = 15,
  pt.alpha = 1,
  line.alpha = 0)
```

features	<i>Feature Points</i>
----------	-----------------------

Description

Get point indices for features, usually for use with [subset_tem](#).

Usage

```
features(..., tem_id = c("frl", "dlib70"))
```

Arguments

...	a vector of feature names (see Details)
tem_id	template ID (currently only works for frl and dlib70)

Details

Available features for the frl template are: "gmm", "oval", "face", "mouth", "nose", "eyes", "brows", "left_eye", "right_eye", "left_brow", "right_brow", "ears", "undereyes", "teeth", "smile_lines", "cheek-bones", "philtrum", "chin", "neck", "halo".

Available features for the dlib70 template are: "teeth", "left_eye", "right_eye", "left_brow", "right_brow", "nose", "mouth", "face".

Value

vector of corresponding template indices

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Examples

```
features("mouth")
features("gmm")
features("nose", tem_id = "dlib70")
```

`get_info`*Get Information*

Description

Get Information

Usage

```
get_info(stimuli, ..., .rownames = "id")
```

Arguments

<code>stimuli</code>	list of stimuli
<code>...</code>	column names to return
<code>.rownames</code>	whether to return a table with no rownames (NULL), rownames from the list item names (NA), or as a new column (the column name as a string)

Value

a data frame or vector of the info

See AlsoOther info: [add_info\(\)](#), [compare\(\)](#), [get_point\(\)](#), [height\(\)](#), [metrics\(\)](#), [rename_stim\(\)](#), [width\(\)](#)**Examples**

```
stimuli <- demo_stim() |>
  add_info(project = "test", gender = c("F", "M"))

get_info(stimuli)
get_info(stimuli, "gender")
```

`get_point`*Get Point Coordinates*

Description

Get a data frame of the x and y coordinates of a template point

Usage

```
get_point(stimuli, pt = 0)
```

Arguments

stimuli	list of stimuli with templates
pt	point(s) to return

Value

data frame of x and y coordinates of the specified point(s) for each stimulus

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Other info: [add_info\(\)](#), [compare\(\)](#), [get_info\(\)](#), [height\(\)](#), [metrics\(\)](#), [rename_stim\(\)](#), [width\(\)](#)

Examples

```
demo_stim() |> get_point(0:1)
```

gglabel

Label with ggplot annotations

Description

Label image using [ggplot2::annotate](#). All arguments are vectorised over the stimuli and values are recycled or truncated if there are fewer or more than stimuli.

Usage

```
gglabel(stimuli, label = TRUE, x = 0.5, y = 0.95, geom = "text", ...)
```

Arguments

stimuli	list of stimuli
label	a vector of the label text(s) or TRUE to use stimlist names
x	x-coordinate for label anchor (left is 0); values ≤ 1 are interpreted as proportions of width
y	y-coordinate for label anchor (bottom is 0); values ≤ 1 are interpreted as proportions of height
geom	the geom to use
...	further arguments to pass to ggplot2::annotate()

Value

stimlist with labelled images

See Also

[label\(\)](#) for a labeller using syntax like [magick::image_annotate](#)

Visualisation functions [as_ggplot\(\)](#), [draw_tem\(\)](#), [label\(\)](#), [mlabel\(\)](#), [plot.stimlist\(\)](#), [plot.stim\(\)](#), [plot_rows\(\)](#), [plot_stim\(\)](#)

Examples

```
stimuli <- demo_stim()

# label with image names
# the default text size in ggplot is tiny
gglabel(stimuli)

# add a watermark
gglabel(stimuli,
        label = "watermark",
        x = 0.5,
        y = 0.5,
        geom = "text",
        size = 30,
        color = "black",
        angle = -30,
        alpha = 0.5)
```

greyscale

Make images greyscale

Description

Make images greyscale

Usage

```
greyscale(stimuli)
```

```
grayscale(stimuli)
```

Arguments

stimuli list of class stimuli

Value

stimlist with new images

See Also

Stimulus manipulation functions [align\(\)](#), [crop_tem\(\)](#), [crop\(\)](#), [horiz_eyes\(\)](#), [image_func\(\)](#), [mask_oval\(\)](#), [mask\(\)](#), [mirror\(\)](#), [pad\(\)](#), [resize\(\)](#), [rotate\(\)](#), [to_size\(\)](#)

Stimulus manipulation functions [align\(\)](#), [crop_tem\(\)](#), [crop\(\)](#), [horiz_eyes\(\)](#), [image_func\(\)](#), [mask_oval\(\)](#), [mask\(\)](#), [mirror\(\)](#), [pad\(\)](#), [resize\(\)](#), [rotate\(\)](#), [to_size\(\)](#)

Examples

```
stimuli <- demo_stim()
grey_stim <- greyscale(stimuli)
plot(grey_stim)
```

height

Image heights

Description

Image heights

Usage

```
height(stimuli, type = c("all", "min", "max", "unique"))
```

Arguments

stimuli	list of stimuli
type	whether to return all heights, min, max, or only unique heights

Value

vector of heights

See Also

Other info: [add_info\(\)](#), [compare\(\)](#), [get_info\(\)](#), [get_point\(\)](#), [metrics\(\)](#), [rename_stim\(\)](#), [width\(\)](#)

Examples

```
demo_stim() |> height()
```

horiz_eyes	<i>Make eyes horizontal</i>
------------	-----------------------------

Description

Rotate each stimulus so the eye points are horizontal.

Usage

```
horiz_eyes(stimuli, left_eye = 0, right_eye = 1, fill = wm_opts("fill"))
```

Arguments

stimuli	list of stimuli
left_eye	The first point to align (defaults to 0)
right_eye	The second point to align (defaults to 1)
fill	background color to pass to rotate, see color_conv()

Value

list of stimuli with rotated tems and/or images

See Also

Stimulus manipulation functions [align\(\)](#), [crop_tem\(\)](#), [crop\(\)](#), [greyscale\(\)](#), [image_func\(\)](#), [mask_oval\(\)](#), [mask\(\)](#), [mirror\(\)](#), [pad\(\)](#), [resize\(\)](#), [rotate\(\)](#), [to_size\(\)](#)

Examples

```
stimuli <- demo_unstandard(1:3)
horiz_eyes(stimuli, fill = "red")
```

image_func	<i>Apply a magick function to each image</i>
------------	--

Description

This is a convenience function for applying magick functions that take an image as the first argument and return an image. It's fully vectorised, so you can set separate argument values for each image.

Usage

```
image_func(stimuli, func, ...)
```

Arguments

stimuli	list of stimuli
func	the function or a string with the short name of the magick function (see image_func_types())
...	arguments to pass to the function

Details

These functions only affect the image, not the template. If a function changes the morphology of the image (e.g., "implode"), the template will not alter in the same way.

Value

list of stimuli with new images

See Also

Stimulus manipulation functions [align\(\)](#), [crop_tem\(\)](#), [crop\(\)](#), [greyscale\(\)](#), [horiz_eyes\(\)](#), [mask_oval\(\)](#), [mask\(\)](#), [mirror\(\)](#), [pad\(\)](#), [resize\(\)](#), [rotate\(\)](#), [to_size\(\)](#)

Examples

```
stimuli <- demo_stim() |> resize(0.5)

# make a photographic negative version
image_func(stimuli, "negate")

# set different argument values for each image
image_func(stimuli, "implode", factor = c(0.2, -0.2))

# other image functions
image_func(stimuli, "blur", 5, 3)
image_func(stimuli, "contrast", sharpen = 1)
image_func(stimuli, "oilpaint", radius = 5)
image_func(stimuli, "colorize", opacity = 50,
           color = c("hotpink", "dodgerblue"))

# load a logo image and superimpose it on each image
logo <- system.file("extdata/logo.png", package = "webmorphR") |>
  magick::image_read() |>
  magick::image_resize(70)

image_func(stimuli, "composite", logo, offset = "+5+10")

# use a self-defined function
testfunc <- function(image) {
  rot <- magick::image_rotate(image, 180)
  c(image, rot) |> magick::image_average()
}
image_func(stimuli, testfunc)
```

image_func_types	<i>Possible functions</i>
------------------	---------------------------

Description

`image_func` can take a named function from the `magick` package, but only functions that return an image that is compatible with the current template (e.g., doesn't change size or shape).

Usage

```
image_func_types()
```

Value

list of compatible function names

Examples

```
image_func_types()
```

label	<i>Label images</i>
-------	---------------------

Description

Defaults to `mlabel()` unless you use arguments specific to `gglabel()`. All arguments are vectorised over the stimuli and values are recycled or truncated if there are fewer or more than stimuli.

Usage

```
label(stimuli, ...)
```

Arguments

stimuli	list of stimuli
...	arguments to pass on to <code>mlabel()</code> or <code>gglabel()</code>

Value

stimlist with labelled images

See Also

`mlabel()`, `gglabel()`

Visualisation functions `as_ggplot()`, `draw_tem()`, `gglabel()`, `mlabel()`, `plot.stimlist()`, `plot.stim()`, `plot_rows()`, `plot_stim()`

Examples

```
stimuli <- demo_stim()

# label with magick::image_annotate
label(stimuli,
      text = c("CHINWE", "GEORGE"),
      gravity = c("north", "south"),
      color = "red")

# label with ggplot2::annotate
label(stimuli,
      label = c("CHINWE", "GEORGE"),
      x = 0.5,
      y = c(0.99, 0.02),
      vjust = c(1, 0),
      size = 18,
      color = "red")
```

loop

Loop

Description

Morph between each image in a list of stimuli, looping back to the start.

Usage

```
loop(stimuli, steps = 10, ...)
```

Arguments

stimuli	list of stimuli to morph between
steps	number of steps from one image to the next
...	arguments to pass to trans()

Value

list of stimuli containing each step of the loop

See Also

WebMorph.org functions [avg\(\)](#), [continuum\(\)](#), [symmetrize\(\)](#), [trans\(\)](#), [webmorph_up\(\)](#)

Examples

```
if (webmorph_up()) {
  # align and crop images
  stimuli <- demo_unstandard(1:5) |>
    align() |> crop_tem()

  loop <- loop(stimuli, 5)

  # create an animated gif
  animate(loop, fps = 10)
}
```

mask

Mask Images with templates

Description

Use template points to define the borders of a mask to apply to the images. The image outside of the mask (or inside, if `reverse = TRUE`) is replaced by the fill color.

Usage

```
mask(
  stimuli,
  mask = "face",
  fill = wm_opts("fill"),
  reverse = FALSE,
  expand = 1,
  tem_id = "frl"
)
```

Arguments

<code>stimuli</code>	list of stimuli
<code>mask</code>	vector of masks or a custom list of template points
<code>fill</code>	color to make the mask, see color_conv()
<code>reverse</code>	logical; whether the mask covers the areas outside (<code>FALSE</code>) or inside (<code>TRUE</code>) the mask
<code>expand</code>	how many pixels to expand the mask (negative numbers contract the mask)
<code>tem_id</code>	template ID to pass on to tem_def() to get built-in mask definitions

Details

For FRL templates, the argument `mask` can be a vector with one or more of the following: `oval`, `face`, `neck`, `ears` (`left_ear`, `right_ear`), `eyes` (`left_eye`, `right_eye`), `brows` (`left_brow`, `right_brow`), `mouth`, `teeth`, `nose`.

For Face++ templates (`fpp83` or `fpp106`), the argument `mask` can be a vector with one or more of the following: `face`, `eyes` (`left_eye`, `right_eye`), `brows` (`left_brow`, `right_brow`), `mouth`, `teeth`, `nose`. Because these templates have no forehead points, "face" is usually disappointing.

Set custom masks using the template points (0-based). View an image with labelled templates using `plot(stim, pt.plot = TRUE, pt.shape = "index")`. Separate points along a line with commas, line segments with semicolons, and mask areas with colons. For example, this would be the custom mask for the eyes in the `fpp83` template:

```
"44,4,56,51,79;79,58,11,25,44:61,67,38,34,40;40,41,17,47,61"
```

If you set `expand = 0`, there is sometimes a thin visible line where multiple components of the mask touch.

Value

list of stimuli with masked images

See Also

Stimulus manipulation functions [align\(\)](#), [crop_tem\(\)](#), [crop\(\)](#), [greyscale\(\)](#), [horiz_eyes\(\)](#), [image_func\(\)](#), [mask_oval\(\)](#), [mirror\(\)](#), [pad\(\)](#), [resize\(\)](#), [rotate\(\)](#), [to_size\(\)](#)

Examples

```
stimuli <- demo_stim()

mask(stimuli,
     mask = c("face", "neck", "ears"),
     fill = "dodgerblue")

mask(stimuli, "face", expand = 30)

# reverse masking masks over the features
stimuli |>
  mask("eyes", "#FFFF00", TRUE) |>
  mask("brows", rgb(0.2, 0.5, 0.5), TRUE) |>
  mask("mouth", "#FF000066", TRUE)

# custom mask (list style)
fpp83_eyes <- list(
  left_eye = list(
    c(44,4,56,51,79),
    c(79,58,11,25,44)
  ),
  right_eye = list(
    c(61,67,38,34,40),
```

```

        c(40, 41, 17, 47, 61)
    )
)

demo_tems("fpp83") |>
  mask(fpp83_eyes, fill = color_conv("dodgerblue", alpha = 0.5))

```

mask_oval

Apply an oval mask to images

Description

Superimpose an oval mask on a set of images.

Usage

```
mask_oval(stimuli, bounds = NULL, fill = wm_opts("fill"), each = TRUE)
```

Arguments

stimuli	list of stimuli
bounds	bounds (t, r, b, l) of oval, calculated from templates if NULL
fill	background color for mask, see color_conv()
each	logical; whether to calculate a mask for each image (default) or just one

Details

If the images have templates and bounds = NULL, the maximum and minimum x and y coordinates for each image will be calculated (or the overall max and min if each = FALSE) and an oval with those dimensions and position will be placed over the face.

If bounds are set to a list of top, right, bottom and left boundaries, these will be used instead of the boundaries derived from templates.

Value

list of stimuli with cropped tems and/or images

See Also

Stimulus manipulation functions [align\(\)](#), [crop_tem\(\)](#), [crop\(\)](#), [greyscale\(\)](#), [horiz_eyes\(\)](#), [image_func\(\)](#), [mask\(\)](#), [mirror\(\)](#), [pad\(\)](#), [resize\(\)](#), [rotate\(\)](#), [to_size\(\)](#)

Examples

```
# remove external template points and crop
stimuli <- demo_stim() |> subset_tem(features("face")) |> crop_tem(25)

# three styles of mask
omask1 <- mask_oval(stimuli) |> label("default")
omask2 <- mask_oval(stimuli, each = FALSE) |> label("each = FALSE")
omask3 <- mask_oval(stimuli, bounds = list(t= 50, r = 30, b = 40, l = 30)) |>
  label("manual bounds")

# visualise masks
c(omask1, omask2, omask3) |> plot(nrow = 2, byrow = FALSE)
```

metrics

Image shape metrics

Description

Get metrics defined by template points.

Usage

```
metrics(stimuli, formula = c(0, 1))
```

Arguments

stimuli	list of stimuli with tems
formula	a vector of two points to measure the distance apart, or a string of the formula for the metric

Details

Reference x and y coordinates by point number like $x[0]$ or $y[188]$. Use any R functions to process the numbers, as well as `pow()` (same as `^()`, for consistency with `webmorph.org`). Remember that 0,0 is the top left for images; e.g., `min(y[0], y[1])` gives your the *higher* of the two pupil y-coordinates.

Value

named vector of the metric

See Also

Other info: [add_info\(\)](#), [compare\(\)](#), [get_info\(\)](#), [get_point\(\)](#), [height\(\)](#), [rename_stim\(\)](#), [width\(\)](#)

Examples

```
stimuli <- demo_stim()

metrics(stimuli, c(0, 1)) # eye-spacing

# face width-to-height ratio
fwh <- "abs(max(x[113],x[112],x[114])-min(x[110],x[111],x[109]))/abs(y[90]-min(y[20],y[25]))"
metrics(stimuli, fwh)
```

 mirror

Mirror templates and images

Description

Use `tem_id` to get the symmetry map for your template. If `tem_id` is omitted, images and templates will be fully reversed (e.g., if point 1 is the left eye in the original image, it will be the right eye in the mirrored image).

Usage

```
mirror(stimuli, tem_id = NULL, axis = "vertical")
```

Arguments

<code>stimuli</code>	list of stimuli
<code>tem_id</code>	template ID to be passed to <code>tem_def</code> (usually "frl" or "fpp106") or NULL
<code>axis</code>	vertical or horizontal axis of mirroring

Value

list of stimuli with mirrored images and templates

See Also

Stimulus manipulation functions [align\(\)](#), [crop_tem\(\)](#), [crop\(\)](#), [greyscale\(\)](#), [horiz_eyes\(\)](#), [image_func\(\)](#), [mask_oval\(\)](#), [mask\(\)](#), [pad\(\)](#), [resize\(\)](#), [rotate\(\)](#), [to_size\(\)](#)

Examples

```
# load an image and mirror it
o <- demo_tems("frl") |> resize(0.5)
m <- mirror(o, "frl")

# visualise the face outline points
c(o, m) |>
  subset_tem(features("face")) |>
  draw_tem(pt.shape = "index", pt.size = 15) |>
  label(c("original", "mirrored"))
```

mlabel	<i>Label with magick annotations</i>
--------	--------------------------------------

Description

Label image using [magick::image_annotate](#). All arguments are vectorised over the stimuli and values are recycled or truncated if there are fewer or more than stimuli. Setting a font, weight, style only works if your imagemagick is compiled with fontconfig support.

Usage

```
mlabel(
  stimuli,
  text = TRUE,
  gravity = "north",
  location = "+0+0",
  degrees = 0,
  size = 0.1,
  font = "sans",
  style = "normal",
  weight = 400,
  kerning = 0,
  decoration = NULL,
  color = "black",
  strokecolor = NULL,
  boxcolor = NULL
)
```

Arguments

stimuli	list of stimuli
text	a vector of the label text(s) or TRUE to use stimlist names
gravity	string with gravity value from <code>magick::gravity_types</code> .
location	geometry string with location relative to gravity
degrees	rotates text around center point
size	font size in pixels or proportion of image width (if < 1.0)
font	string with font family such as "sans", "mono", "serif", "Times", "Helvetica", "Trebuchet", "Georgia", "Palatino" or "Comic Sans".
style	value of <code>magick::style_types()</code> : "Undefined", "Any", "Italic", "Normal", "Oblique"
weight	thickness of the font, 400 is normal and 700 is bold.
kerning	increases or decreases whitespace between letters
decoration	value of <code>magick::decoration_types()</code> : "LineThrough" "None", "Overline", "Underline"

color a vector of the label colour(s)
 strokecolor adds a stroke (border around the text)
 boxcolor adds a background color

Value

stimlist with labelled images

See Also

[gglabel\(\)](#) for a labeller using syntax like [ggplot2::annotate\(\)](#)

Visualisation functions [as_ggplot\(\)](#), [draw_tem\(\)](#), [gglabel\(\)](#), [label\(\)](#), [plot.stimlist\(\)](#), [plot.stim\(\)](#), [plot_rows\(\)](#), [plot_stim\(\)](#)

Examples

```
stimuli <- demo_stim()
mlabel(stimuli,
  text = c("CHINWE", "GEORGE"),
  gravity = c("north", "south"),
  color = "red")
```

 pad

Pad images

Description

Add padding to sides of stimuli. This is a convenience function to calculate offsets for [crop\(\)](#).

Usage

```
pad(stimuli, top = 10, right = top, bottom = top, left = right, ...)
```

Arguments

stimuli list of stimuli
 top, right, bottom, left
 number of pixels or proportion (<1) to pad each side
 ... additional arguments to pass to [crop\(\)](#)

Details

The value for top is copied to bottom and right, and the value for right is copied to left, so setting only top produces a consistent border, while setting just top and right sets different borders for top-bottom and right-left. (This convention will be familiar if you use CSS.)

Padding size values are interpreted as a proportion of width or height if less than 1.

Value

list of stimuli

See Also

Stimulus manipulation functions [align\(\)](#), [crop_tem\(\)](#), [crop\(\)](#), [greyscale\(\)](#), [horiz_eyes\(\)](#), [image_func\(\)](#), [mask_oval\(\)](#), [mask\(\)](#), [mirror\(\)](#), [resize\(\)](#), [rotate\(\)](#), [to_size\(\)](#)

Examples

```
stimuli <- demo_stim()

# default 10-pixel padding
pad(stimuli, fill = "dodgerblue")

# change pad width and set fill
pad(stimuli, 2, fill = "dodgerblue")

# set top border to 10% height
# different colour for each image
pad(stimuli, 0.1, 1, 1, 1,
     fill = c("hotpink", "dodgerblue"))
```

patch

Patch colour

Description

Get the median (or mean or user-defined function) colour value of a specified patch of pixels on an image. This is useful for matching background colours.

Usage

```
patch(
  stimuli,
  width = 10,
  height = 10,
  x_off = 0,
  y_off = 0,
  color = c("hex", "rgb"),
  func = stats::median
)
```

Arguments

stimuli	list of stimuli
width, height	dimensions of the patch in pixels, if ≤ 1 , interpreted as proportions of width or height
x_off, y_off	offset in pixels or proportion (< 1)
color	The type of color to return (hex, rgb)
func	The function to apply to an array of L*ab color values to determine the central colour (defaults to median, but mean, min, or max can also be useful)

Details

The colour values of each pixel in the patch are converted to CIE-Lab values before using the func to calculate the central tendency of the L (lightness), a (red-green axis) and b (blue-yellow axis); see `col2lab()` and `lab2rgb()` for more details.

This excludes transparent pixels, and returns "transparent" if all pixels in the patch are transparent.

Value

a vector of hex or rgba color values

Examples

```
stimuli <- demo_stim()

# get colour from the upper left corner
patch(stimuli)

# get median colour from centre .1 width pixels
patch(stimuli, width = .1, height = .1,
      x_off = .45, y_off = .45)

# get mean rgb colour from full image
patch(stimuli, width = 1, height = 1,
      color = "rgb", func = mean)
```

plot_rows

Plot in rows

Description

Plot in rows

Usage

```
plot_rows(  
  ...,  
  top_label = NULL,  
  maxwidth = wm_opts("plot.maxwidth"),  
  maxheight = wm_opts("plot.maxheight")  
)
```

Arguments

... stimlists (optionally named) and any arguments to pass on to [label](#)

top_label logical; whether to plot row labels above the row (TRUE) or inside (FALSE), if NULL, then TRUE if stimlists are named

maxwidth, maxheight maximum width and height of each row in pixels

Value

stimlist with plot

See Also

Visualisation functions [as_ggplot\(\)](#), [draw_tem\(\)](#), [gglabel\(\)](#), [label\(\)](#), [mlabel\(\)](#), [plot.stimlist\(\)](#), [plot.stim\(\)](#), [plot_stim\(\)](#)

Examples

```
s <- demo_unstandard()  
plot_rows(  
  female = s[1:3],  
  male = s[6:8],  
  maxwidth = 600  
)
```

plot_stim

Plot stimuli

Description

Show all the stimuli in a grid. You can use [plot\(\)](#) as an alias.

Usage

```
plot_stim(
  stimuli,
  nrow = NULL,
  ncol = NULL,
  byrow = TRUE,
  padding = 10,
  external_pad = TRUE,
  fill = wm_opts("fill"),
  maxwidth = wm_opts("plot.maxwidth"),
  maxheight = wm_opts("plot.maxheight")
)
```

Arguments

stimuli	list of class stimlist
nrow	number of rows
ncol	number of columns
byrow	fill grid by rows (first ncol images in the first row); if FALSE, fills by columns (first nrow images in the first column)
padding	around each image in pixels
external_pad	whether to include external padding
fill	background color, see color_conv()
maxwidth, maxheight	maximum width and height of grid in pixels

Value

stimlist with the plot image (no templates)

See Also

Visualisation functions [as_ggplot\(\)](#), [draw_tem\(\)](#), [gglabel\(\)](#), [label\(\)](#), [mlabel\(\)](#), [plot.stimlist\(\)](#), [plot.stim\(\)](#), [plot_rows\(\)](#)

Examples

```
stimuli <- demo_stim() |> resize(0.5)
plot_stim(stimuli)

# default padding is 10px internal and external
plot(stimuli, fill = "dodgerblue")
plot(stimuli, external_pad = 0, fill = "dodgerblue")
plot(stimuli, padding = 0, fill = "dodgerblue")

# make 8 numbered images
n <- blank(8, color = grDevices::cm.colors(8)) |>
```

```
    label(1:8, gravity = "center", size = 50)

# 2 rows, allocating by row
plot(n, nrow = 2)

# 2 rows, allocating by column
plot(n, nrow = 2, byrow = FALSE)
```

read_stim

Read stimuli

Description

Read images and templates from a directory.

Usage

```
read_stim(path, pattern = NULL, breaks = "/")
```

Arguments

path	Path to directory containing image and/or template files (or a single file path)
pattern	Vector of patterns to use to search for files, or a vector of image indices (e.g., 1:4 selects the first 4 images and their templates if they exist)
breaks	a vector of characters used to determine the stimulus names from the file names

Value

a list of stimuli

See Also

Stimulus creating functions [animate\(\)](#), [as_stimlist\(\)](#), [blank\(\)](#), [new_stimlist\(\)](#), [new_stim\(\)](#), [read_img\(\)](#), [read_tem\(\)](#), [write_stim\(\)](#)

Examples

```
path <- system.file("extdata/test", package = "webmorphR")

# read in all images and templates in a directory
stimuli <- read_stim(path)

# read in just images and templates with "m_"
m_stimuli <- read_stim(path, "m_")
```

remove_tem	<i>Remove templates</i>
------------	-------------------------

Description

Remove templates

Usage

```
remove_tem(stimuli)
```

Arguments

stimuli list of stimuli

Value

list of stimuli

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [get_point\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Examples

```
demo_stim() |> remove_tem()
```

rename_stim	<i>Set stimulus names in a stimlist</i>
-------------	---

Description

Set stimulus names in a stimlist

Usage

```
rename_stim(
  stimuli,
  new_names = NULL,
  prefix = "",
  suffix = "",
  pattern = NULL,
  replacement = NULL,
  ...
)
```

Arguments

stimuli	A list of stimuli
new_names	Vector of new names - must be the same length as the stimlist
prefix	String to prefix to each name
suffix	String to append to each name
pattern	Pattern for gsub
replacement	Replacement for gsub
...	Additional arguments to pass on to <code>base::gsub()</code>

Value

A list of stimuli with the new names

See Also

Other info: [add_info\(\)](#), [compare\(\)](#), [get_info\(\)](#), [get_point\(\)](#), [height\(\)](#), [metrics\(\)](#), [width\(\)](#)

Examples

```
demo_stim() |>
  rename_stim(prefix = "new_") |>
  names()
```

require_tems

Require templates

Description

Checks a list of stimuli for templates and omits images without a template. If `all_same = TRUE`, checks that all the templates are the same type. Errors if no images have a template or not all templates are the same (when `all_same == TRUE`).

Usage

```
require_tems(stimuli, all_same = FALSE)
```

Arguments

stimuli	list of stimuli
all_same	logical; whether all images should have the same template

Value

list of stimuli with tems

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Examples

```
stimuli <- demo_stim()
have_tems <- require_tems(stimuli)

## Not run:
# produces an error because no tems
no_tems <- stimuli |> remove_tem()
require_tems(no_tems)

# warns that some images were removed
mix_tems <- c(stimuli, no_tems)
have_tems <- require_tems(mix_tems)

# produces an error because tems are different
demo_tems() |> require_tems(all_same = TRUE)

## End(Not run)
```

 resize

Resize stimuli

Description

Resize images and templates to the specified width and height.

Usage

```
resize(stimuli, width = NULL, height = NULL)
```

Arguments

`stimuli` list of stimuli
`width, height` new dimensions (in pixels or percent if < 10)

Value

list of stimuli with resized tems and/or images

See Also

Stimulus manipulation functions [align\(\)](#), [crop_tem\(\)](#), [crop\(\)](#), [greyscale\(\)](#), [horiz_eyes\(\)](#), [image_func\(\)](#), [mask_oval\(\)](#), [mask\(\)](#), [mirror\(\)](#), [pad\(\)](#), [rotate\(\)](#), [to_size\(\)](#)

Examples

```
stimuli <- demo_stim()

# set width to proportion, height proportional
resize(stimuli, .5) |> draw_tem()

# set width and height separately by pixels
resize(stimuli, 400, 250) |> draw_tem()
```

rotate	<i>Rotate templates and images</i>
--------	------------------------------------

Description

Rotate templates and images

Usage

```
rotate(
  stimuli,
  degrees = 0,
  fill = wm_opts("fill"),
  keep_size = TRUE,
  origin = "image"
)
```

Arguments

stimuli	list of stimuli
degrees	degrees to rotate
fill	background color, see color_conv()
keep_size	whether to keep the original size or expand images to the new rotated size
origin	The origin of the rotation. Options are: "image" will rotate around the image center. "tem" will rotate around the average of all template coordinates. A vector of 1 or more point indices (0-based) will rotate around their average position.

Value

list of stimuli with rotated tems and/or images

See Also

Stimulus manipulation functions [align\(\)](#), [crop_tem\(\)](#), [crop\(\)](#), [greyscale\(\)](#), [horiz_eyes\(\)](#), [image_func\(\)](#), [mask_oval\(\)](#), [mask\(\)](#), [mirror\(\)](#), [pad\(\)](#), [resize\(\)](#), [to_size\(\)](#)

Examples

```

stimuli <- demo_stim() |> resize(0.5)

rotate(stimuli, 45, fill = "dodgerblue")
rotate(stimuli, 45, fill = "dodgerblue", keep_size = FALSE)

# if images are not in the centre of the image,
# try setting the origin to tem or specific point(s)
offset <- stimuli[1] |>
  draw_tem() |>
  pad(0, 250, 0, 0, fill = "dodgerblue")

rotate(offset, 45, origin = "image", fill = "pink")
rotate(offset, 45, origin = "tem", fill = "pink")

# rotate around point 0 (left eye)
offset |> crop_tem() |> rep(8) |>
  rotate(seq(0, 325, 45), origin = 0, fill = "pink") |>
  animate(fps = 5)

```

same_tems

Check All Templates are the Same

Description

Check All Templates are the Same

Usage

```
same_tems(stimuli)
```

Arguments

stimuli list of stimuli

Value

logical

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Examples

```
stim <- demo_stim()
stim2 <- subset_tem(stim, features("gmm"))

same_tems(stim)

c(stim, stim2) |> same_tems()
```

social_media_size	<i>Social Media Image Sizes</i>
-------------------	---------------------------------

Description

A convenience function for getting recommended dimensions for images on social media sites.

Usage

```
social_media_size(platform = c("twitter", "instagram"), type = "default")
```

Arguments

platform	currently only "twitter"
type	which type of image

Details

Twitter:

link: Image from a Tweet with shared link one: Tweet sharing a single image (default) two: Tweet sharing two images three_left: Tweet sharing three images, Left image three_tight Tweet sharing three images, Right images four: Tweet sharing four images

Instagram:

feed_large: (default) feed_small: stories_large: stories_small:

Value

named vector of width and height in pixels

Examples

```
social_media_size("twitter", "link")
social_media_size("twitter", "one")
social_media_size("twitter", "two")
```

`squash_tem`*Squash Template Points*

Description

Move template points that are outside the image boundaries (e.g., negative values or larger than image width or height) to the borders of the image.

Usage

```
squash_tem(stimuli)
```

Arguments

`stimuli` list of stimuli

Value

list of stimuli

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Examples

```
nosquash <- demo_stim(1) |>
  crop(0.4, 0.5)

squash <- demo_stim(1) |>
  crop(0.4, 0.5) |>
  squash_tem()

# add padding and visualise templates
c(nosquash, squash) |>
  pad(50) |>
  draw_tem(pt.size = 5)
```

subset_tem	<i>Subset template points</i>
------------	-------------------------------

Description

Keep or delete specified template points. Points will be renumbered and line definitions will be updated. If all points in a line are deleted, the line will be removed. POint indexing is 0-based, so the first two points (usually the pupils) are 0 and 1.

Usage

```
subset_tem(stimuli, ..., keep = TRUE)
```

Arguments

stimuli	list of stimuli
...	vectors of points to keep or delete
keep	logical; whether to keep or delete the points

Value

stimlist with altered templates

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [tem_def\(\)](#), [viz_tem_def\(\)](#)

Examples

```
# keep just the first two points
demo_stim(1) |>
  subset_tem(0:1) |>
  draw_tem(pt.size = 10)

# remove the last 10 points
# (produces the 179-point Perception Lab template)
demo_stim(1) |>
  subset_tem(179:188, keep = FALSE) |>
  draw_tem()

# use features() to keep only points from a pre-defined set
# "gmm" is points used for geometric morphometrics
demo_stim(1) |>
  subset_tem(features("gmm")) |>
  draw_tem()
```

`symmetrize`*Symmetrize Images*

Description

Use webmorph.org to make faces symmetric in shape and/or colour.

Usage

```
symmetrize(stimuli, shape = 1, color = 1, tem_id = "frl", ...)
```

```
symmetrise(stimuli, shape = 1, color = 1, tem_id = "frl", ...)
```

Arguments

<code>stimuli</code>	list of stimuli
<code>shape, color</code>	amount of symmetry (0 for none, 1.0 for perfect)
<code>tem_id</code>	template ID to be passed to <code>tem_def()</code> (usually "frl" or "fpp106")
<code>...</code>	Additional arguments to pass to <code>trans()</code>

Value

list of stimuli with symmetrised images and templates

See Also

WebMorph.org functions `avg()`, `continuum()`, `loop()`, `trans()`, `webmorph_up()`

Examples

```
if (webmorph_up()) {
  stimuli <- demo_stim(1)

  sym_both <- symmetrize(stimuli)
  sym_shape <- symmetrize(stimuli, color = 0)
  sym_color <- symmetrize(stimuli, shape = 0)
  sym_anti <- symmetrize(stimuli, shape = -1.0, color = 0)
}
```

tems_to_array	<i>Convert stimuli to array for geomorph</i>
---------------	--

Description

Convert stimuli to array for geomorph

Usage

```
tems_to_array(stimuli)
```

Arguments

stimuli list of stimuli

Value

3D array

Examples

```
data <- demo_stim() |> tems_to_array()
dim(data)
```

tem_def	<i>Get template definition</i>
---------	--------------------------------

Description

Template definitions are lists that contain information about templates that are needed to do things like symmetrising and masking images. This function is mostly used internally.

Usage

```
tem_def(tem_id = "fr1", path = NULL)
```

Arguments

tem_id the name of a built-in template (fr1, fpp106, fpp83, dlib70, or dlib7) or a numeric ID of a template to retrieve from webmorph.org

path path of local tem definition file

Details

If you have defined a custom template on webmorph.org, you can get its function definition by ID. You can see the ID numbers next to the templates available to you under the *Template > Current Template* menu.

Value

list with template definition

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [viz_tem_def\(\)](#)

Examples

```
fpp106 <- tem_def("fpp106")
fpp106$lines |> str()
```

```
fpp83 <- tem_def("fpp83")
fpp83$mask |> str()
```

```
fr1 <- tem_def("fr1")
fr1$points[1:10, ]
viz_tem_def(fr1, pt.size = 10, line.size = 5)
```

to_size

Resize and crop/pad images to a specified size

Description

Resize and crop/pad images to a specified size

Usage

```
to_size(
  stimuli,
  width = NULL,
  height = NULL,
  fill = wm_opts("fill"),
  crop = FALSE,
  keep_rels = FALSE
)
```


Arguments

stimuli	list of stimuli
width	the target width (if null, the maximum stimulus width is used)
height	the target height (if null, the maximum stimulus height is used)
fill	background color if cropping goes outside the original image, see color_conv()
crop	whether to crop or pad images to make them the specified size
keep_rels	whether to keep the size relationships between images in the set, or make all the maximum size

Value

list of stimuli with cropped tems and/or images

See Also

Stimulus manipulation functions [align\(\)](#), [crop_tem\(\)](#), [crop\(\)](#), [greyscale\(\)](#), [horiz_eyes\(\)](#), [image_func\(\)](#), [mask_oval\(\)](#), [mask\(\)](#), [mirror\(\)](#), [pad\(\)](#), [resize\(\)](#), [rotate\(\)](#)

Examples

```
# images with different aspect ratios and sizes
stimuli <- demo_unstandard(c(1:4, 6:9))

to_size(stimuli, 200, 200, fill = "dodgerblue")
```

trans

Transform Images

Description

Transform a base image in shape, color, and/or texture by the differences between two images.

Usage

```
trans(
  trans_img = NULL,
  from_img = NULL,
  to_img = NULL,
  shape = 0,
  color = 0,
  texture = 0,
  outname = NULL,
  norm = c("none", "twopoint", "rigid"),
  normpoint = 0:1,
```

```

    sample_contours = TRUE,
    warp = c("multiscale", "linear", "multiscalerb")
)

```

Arguments

trans_img	list of stimuli to transform
from_img	negative transform dimension endpoint (0% image)
to_img	positive transform dimension endpoint (100% image)
shape, color, texture	amount to change along the vector defined by from_img and to_img (can range from -3 to +3)
outname	name to save each image as
norm	how to normalise the images; see Details
normpoint	points for twopoint normalisation
sample_contours	whether to sample contours or just points
warp	warp type

Details

Normalisation options:

- none: averages will have all coordinates as the mathematical average of the coordinates in the component templates
- twopoint: all images are first aligned to the 2 alignment points designated in normpoint. Their position is set to their position in the first image in stimuli
- rigid: procrustes aligns all images to the position of the first image in stimuli

Sample contours:

This interpolates more control points along the lines. This can improve the accuracy of averages and transforms. If you see a “feathery” appearance along lines that have many, close-together points, try turning this off.

Warp types:

- multiscale: Implements multi-scale affine interpolation for image warping. This is the default, with a good balance between speed and accuracy
- linear: Implements triangulated linear interpolation for image warping. Linear warping is least accurate, often resulting in image artifacts, but is very fast.
- multiscalerb: Implements multi-scale rigid body interpolation for image warping. This decreases image artifacts in some circumstances, but is much slower.

Value

list of stimuli with transformed images and templates

See Also

WebMorph.org functions [avg\(\)](#), [continuum\(\)](#), [loop\(\)](#), [symmetrize\(\)](#), [webmorph_up\(\)](#)

Examples

```
if (webmorph_up()) {
  stimuli <- demo_stim()
  sexdim <- trans(stimuli, stimuli$f_multi, stimuli$m_multi,
                 shape = c(fem = -0.5, masc = 0.5))

  sexdim |> draw_tem() |> label()
}
```

 viz_tem_def

Visualise a template definition

Description

Visualise a template definition

Usage

```
viz_tem_def(tem_def, ...)
```

Arguments

tem_def	the template definition; usually from tem_def()
...	further arguments to pass to draw_tem() ; pt.size and line.size often need to be adjusted

Value

a stimlist with a blank image and the template drawn on it

See Also

Template functions [auto_delin\(\)](#), [average_tem\(\)](#), [centroid\(\)](#), [change_lines\(\)](#), [delin\(\)](#), [draw_tem\(\)](#), [features\(\)](#), [get_point\(\)](#), [remove_tem\(\)](#), [require_tems\(\)](#), [same_tems\(\)](#), [squash_tem\(\)](#), [subset_tem\(\)](#), [tem_def\(\)](#)

Examples

```
dlib70 <- tem_def("dlib70")
viz_tem_def(dlib70, pt.size = 5, line.size = 3)
```

webmorph_up	<i>Check if webmorph.org is available</i>
-------------	---

Description

Check if webmorph.org is available

Usage

```
webmorph_up()
```

See Also

WebMorph.org functions [avg\(\)](#), [continuum\(\)](#), [loop\(\)](#), [symmetrize\(\)](#), [trans\(\)](#)

Examples

```
webmorph_up()
```

width	<i>Image widths</i>
-------	---------------------

Description

Image widths

Usage

```
width(stimuli, type = c("all", "min", "max", "unique"))
```

Arguments

stimuli	list of stimuli
type	whether to return all widths, min, max, or only unique widths

Value

vector of widths

See Also

Other info: [add_info\(\)](#), [compare\(\)](#), [get_info\(\)](#), [get_point\(\)](#), [height\(\)](#), [metrics\(\)](#), [rename_stim\(\)](#)

Examples

```
demo_stim() |> width()
```

`wm_opts`*Set/get global webmorph options*

Description

See [wm_opts_defaults\(\)](#) for explanations of the default options.

Usage

```
wm_opts(...)
```

Arguments

... One of four: (1) nothing, then returns all options as a list; (2) a name of an option element, then returns its value; (3) a name-value pair which sets the corresponding option to the new value (and returns nothing), (4) a list with option-value pairs which sets all the corresponding arguments.

Value

a list of options, values of an option, or nothing

See Also

[wm_opts_defaults\(\)](#)

Examples

```
wm_opts() # see all options

wm_opts("verbose") # see value of webmorph.verbose

## Not run:
# set value of webmorph.verbose
wm_opts(verbose = FALSE)

# set multiple options
opts <- list(fill = "black",
             pt.color = "white",
             line.color = "red")
wm_opts(opts)

## End(Not run)
```

wm_opts_defaults *WebmorphR default options*

Description

Options set on load (unless they were already set by `.Renviron`)

- `overwrite` ("ask"): Whether to overwrite images saved with `write_stim()` when in interactive mode; possible values are "ask" (ask if filenames exist), `TRUE` (always overwrite), and `FALSE` (never overwrite)
- `fill` ("white"): the colour to use to fill image backgrounds
- `pt.color` ("green"): the colour to use for points in `draw_tem()`
- `line.color` ("blue"): the colour to use for lines in `draw_tem()`
- `plot` ("inline"): whether to plot images inline in R markdown documents (set to any other value to just view them in the viewer)
- `plot.maxwidth` (2400): The maximum width of images created by `plot()`
- `plot.maxheight` (2400): The maximum height of images created by `plot()`
- `verbose` (`TRUE`): Whether to produce verbose output and progress bars for long functions like `auto_delin()`, `avg()` or `trans()`
- `server` ("https://webmorph.org"): The server to use for webmorph functions like `avg()` and `trans()`; do not change unless you've set up a local server
- `connection` (`stdin()`): use internally for testing interactive functions; do not change

Usage

```
wm_opts_defaults()
```

Value

a list of default options

See Also

[wm_opts\(\)](#)

Examples

```
wm_opts_defaults() |> str() # view defaults

## Not run:
# reset all options to default
wm_opts_defaults() |> wm_opts()

## End(Not run)
```

write_stim	<i>Write images and templates to files</i>
------------	--

Description

Write images and templates to files

Usage

```
write_stim(
  stimuli,
  dir = ".",
  names = NULL,
  format = "png",
  ...,
  overwrite = wm_opts("overwrite")
)
```

Arguments

stimuli	list of stimuli
dir	Directory to save to
names	A vector of stimulus names or NULL to use names from the stimuli list
format	output format such as "png", "jpeg", "gif"; is overridden if names end in .png, .jpg, or .gif
...	other arguments to pass to magick::image_write , such as quality (for jpegs)
overwrite	whether to overwrite existing files (TRUE/FALSE) or "ask" (only in interactive mode)

Value

list of saved paths

See Also

Stimulus creating functions [animate\(\)](#), [as_stimlist\(\)](#), [blank\(\)](#), [new_stimlist\(\)](#), [new_stim\(\)](#), [read_img\(\)](#), [read_stim\(\)](#), [read_tem\(\)](#)

Examples

```
## Not run:
# write demo stim as jpegs to directory ./test_faces
demo_stim() |>
  write_stim("test_faces", format = "jpg")

## End(Not run)
```

write_tps	<i>Create a TPS file from a stimlist</i>
-----------	--

Description

Create a TPS file from a stimlist

Usage

```
write_tps(stimuli, path_to_tps = NULL)
```

Arguments

stimuli	list of stimuli
path_to_tps	optional filename to save TPS file

Value

text of tps file

Examples

```
# set path_to_tps to save to a file
tps <- demo_stim() |>
write_tps()
```


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