Package 'rphylopic'

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Title Get Silhouettes of Organisms from PhyloPic

```
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```

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Description

Specify existing images, taxonomic names, or PhyloPic uuids to add PhyloPic silhouettes as a separate layer to an existing ggplot plot.

Usage

```
add_phylopic(
  img = NULL,
  name = NULL,
  uuid = NULL,
  filter = NULL,
  x,
  y,
  ysize = deprecated(),
  height = NA,
  width = NA,
  alpha = 1,
  color = NA,
```

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```
fill = "black",
horizontal = FALSE,
vertical = FALSE,
angle = 0,
hjust = 0.5,
vjust = 0.5,
remove_background = TRUE,
verbose = FALSE
)
```

Arguments

uuid

img A Picture or png array object, e.g., from using get_phylopic().

name character. A taxonomic name to be passed to get_uuid().

character. A valid uuid for a PhyloPic silhouette (such as that returned by

get_uuid() or pick_phylopic()).

filter character. Filter by usage license if name is defined. Use "by" to limit results

to images which do not require attribution, "nc" for images which allows commercial usage, and "sa" for images without a ShareAlike clause. The user can

also combine these filters as a vector.

x numeric. x value of the silhouette center.

y numeric. y value of the silhouette center.

ysize [Deprecated] use the height or width argument instead.

height numeric. Height of the silhouette in coordinate space. If "NA", the default, and

width is "NA", the silhouette will be as large as fits in the plot area. If "NA" and width is specified, the height is determined by the aspect ratio of the original

image. One or both of height and width must be "NA".

width numeric. Width of the silhouette in coordinate space. If "NA", the default, and

height is "NA", the silhouette will be as large as fits in the plot area. If "NA", the default, and height is specified, the width is determined by the aspect ratio

of the original image. One or both of height and width must be "NA".

alpha numeric. A value between 0 and 1, specifying the opacity of the silhouette (0 is

fully transparent, 1 is fully opaque).

color character. Color of silhouette outline. If "original" or NA is specified, the

original color of the silhouette outline will be used (usually the same as "trans-

parent"). To remove the outline, you can set this to "transparent".

fill character. Color of silhouette. If "original" is specified, the original color of

the silhouette will be used (usually the same as "black"). If color is specified and fill is NA, color will be used as the fill color (for backwards compatibil-

ity). To remove the fill, you can set this to "transparent".

horizontal logical. Should the silhouette be flipped horizontally?

vertical logical. Should the silhouette be flipped vertically?

angle numeric. The number of degrees to rotate the silhouette clockwise. The default

is no rotation.

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hjust numeric. A numeric vector between 0 and 1 specifying horizontal justification (left = 0, center = 0.5, right = 1). Note that, due to the enforcement of the silhouette's aspect ratio, there may be unexpected behavior due to interactions between the aspect ratio of the plot and the aspect ratio of the silhouette.

numeric. A numeric vector between 0 and 1 specifying vertical justification (top = 1, middle = 0.5, bottom = 0). Note that, due to the enforcement of the silhouette's aspect ratio, there may be unexpected behavior due to interactions between the aspect ratio of the plot and the aspect ratio of the silhouette.

remove_background

logical. Should any white background be removed from the silhouette(s)? See

recolor_phylopic() for details.

logical. Should the attribution information for the used silhouette(s) be printed

to the console (see get_attribution())?

Details

vjust

verbose

One (and only one) of img, name, or unid must be specified. Use parameters x, y, and ysize to place the silhouette at a specified position on the plot. The aspect ratio of the silhouette will always be maintained.

x and/or y may be vectors of numeric values if multiple silhouettes should be plotted at once. In this case, any other arguments (except for remove_background) may also be vectors of values, which will be recycled as necessary.

When specifying a horizontal and/or vertical flip **and** a rotation, the flip(s) will always occur first. If you would like to customize this behavior, you can flip and/or rotate the image within your own workflow using flip_phylopic() and rotate_phylopic().

Note that png array objects can only be rotated by multiples of 90 degrees. Also, outline colors do not currently work for png array objects.

Examples

```
# Put a silhouette behind a plot based on a taxonomic name
library(ggplot2)
ggplot(iris) +
  add_phylopic(x = 6.1, y = 3.2, name = "Iris", alpha = 0.2) +
  geom_point(aes(x = Sepal.Length, y = Sepal.Width))
# Put a silhouette in several places based on UUID
posx <- runif(10, 0, 10)
posy <- runif(10, 0, 10)
heights <- runif(10, 0.4, 2)
angle <- runif(10, 0, 360)
hor <- sample(c(TRUE, FALSE), 10, TRUE)
ver <- sample(c(TRUE, FALSE), 10, TRUE)</pre>
fills <- sample(c("black", "darkorange", "grey42", "white"), 10,
  replace = TRUE)
alpha <- runif(10, 0.3, 1)
p <- ggplot(data.frame(cat.x = posx, cat.y = posy), aes(cat.x, cat.y)) +</pre>
```

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add_phylopic_base

Add PhyloPics to a base R plot

Description

Specify existing images, taxonomic names, or PhyloPic unids to add PhyloPic silhouettes on top of an existing base R plot (like points()).

Usage

```
add_phylopic_base(
  img = NULL,
  name = NULL,
  uuid = NULL,
  filter = NULL,
  x = NULL
  y = NULL,
  ysize = deprecated(),
  height = NULL,
 width = NULL,
  alpha = 1,
  color = NA,
  fill = "black",
  horizontal = FALSE,
  vertical = FALSE,
  angle = 0,
  hjust = 0.5,
  vjust = 0.5,
  remove_background = TRUE,
  verbose = FALSE
)
```

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| filter | character. Filter by usage license if name is defined. Use "by" to limit results to images which do not require attribution, "nc" for images which allows commercial usage, and "sa" for images without a ShareAlike clause. The user can also combine these filters as a vector. | |
|-------------------|--|--|
| x | numeric. x value of the silhouette center. If "NULL", the default, the mean value of the x -axis is used. | |
| У | numeric. y value of the silhouette center. If "NULL", the default, the mean value of the y-axis is used. | |
| ysize | [Deprecated] use the height or width argument instead. | |
| height | numeric. Height of the silhouette in coordinate space. If "NULL", the default, and width is also "NULL", the silhouette will be as large as fits in the plot area. If "NULL" and width is specified, the height is determined by the aspect ratio of the original image. One or both of height and width must be "NULL". | |
| width | numeric. Width of the silhouette in coordinate space. If "NULL", the default, and height is also "NULL", the silhouette will be as large as fits in the plot area. If "NULL" and height is specified, the width is determined by the aspect ratio of the original image. One or both of height and width must be "NULL". | |
| alpha | numeric. A value between 0 and 1, specifying the opacity of the silhouette (0 is fully transparent, 1 is fully opaque). | |
| color | character. Color of silhouette outline. If "original" or NA is specified, the original color of the silhouette outline will be used (usually the same as "transparent"). To remove the outline, you can set this to "transparent". | |
| fill | character. Color of silhouette. If "original" is specified, the original color of the silhouette will be used (usually the same as "black"). If color is specified and fill is NA, color will be used as the fill color (for backwards compatibility). To remove the fill, you can set this to "transparent". | |
| horizontal | logical. Should the silhouette be flipped horizontally? | |
| vertical | logical. Should the silhouette be flipped vertically? | |
| angle | numeric. The number of degrees to rotate the silhouette clockwise. The default is no rotation. | |
| hjust | numeric. A numeric vector between 0 and 1 specifying horizontal justification (left = 0 , center = 0.5 , right = 1). | |
| vjust | numeric. A numeric vector between 0 and 1 specifying vertical justification (top = 1, middle = 0.5 , bottom = 0). | |
| remove_background | | |
| | logical. Should any white background be removed from the silhouette(s)? See recolor_phylopic() for details. | |
| verbose | logical. Should the attribution information for the used silhouette(s) be printed to the console (see <pre>get_attribution()</pre> ? | |

Details

One (and only one) of img, name, or unid must be specified. Use parameters x, y, hjust, and vjust to place the silhouette at a specified position on the plot. If height and width are both unspecified,

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then the silhouette will be plotted to the full height and/or width of the plot. The aspect ratio of Picture objects will always be maintained (even when a plot is resized). However, if the plot is resized after plotting a silhouette, the absolute size and/or position of the silhouette may change.

Any argument (except for remove_background) may be a vector of values if multiple silhouettes should be plotted. In this case, all other arguments may also be vectors of values, which will be recycled as necessary to the length of the longest vector argument.

When specifying a horizontal and/or vertical flip **and** a rotation, the flip(s) will always occur first. If you would like to customize this behavior, you can flip and/or rotate the image within your own workflow using flip_phylopic() and rotate_phylopic().

Note that png array objects can only be rotated by multiples of 90 degrees. Also, outline colors do not currently work for png array objects.

Examples

```
## Not run:
# single image
plot(1, 1, type = "n", main = "A cat")
add_phylopic_base(uuid = "23cd6aa4-9587-4a2e-8e26-de42885004c9",
                  x = 1, y = 1, height = 0.4)
# lots of images using a uuid
posx <- runif(10, 0, 1)
posy <- runif(10, 0, 1)
size <- runif(10, 0.1, 0.3)
angle <- runif(10, 0, 360)
hor <- sample(c(TRUE, FALSE), 10, TRUE)</pre>
ver <- sample(c(TRUE, FALSE), 10, TRUE)</pre>
fills <- sample(c("black", "darkorange", "grey42", "white"), 10,
               replace = TRUE)
plot(posx, posy, type = "n", main = "A cat herd")
add_phylopic_base(uuid = "23cd6aa4-9587-4a2e-8e26-de42885004c9",
                  x = posx, y = posy, height = size,
                  fill = fills, angle = angle,
                  horizontal = hor, vertical = ver)
# Example using a cat background
cat <- get_phylopic("23cd6aa4-9587-4a2e-8e26-de42885004c9")</pre>
# setup plot area
plot(posx, posy, type = "n", main = "A cat herd, on top of a cat",
     xlim = c(0, 1), ylim = c(0, 1)
# plot background cat
add_phylopic_base(img = cat, alpha = 0.2)
# overlay smaller cats
add_phylopic_base(img = cat, x = posx, y = posy, height = size, alpha = 0.8)
## End(Not run)
```

add_phylopic_legend Add a PhyloPic legend for a base R plot

Description

Specify existing images, taxonomic names, or PhyloPic unids to add PhyloPic silhouettes as a legend to an existing base R plot (like legend()).

Usage

```
add_phylopic_legend(
    x,
    y = NULL,
    legend,
    img = NULL,
    name = NULL,
    uuid = NULL,
    ysize = deprecated(),
    height = NULL,
    color = NA,
    fill = "black",
    ...
)
```

| X | numeric or character. Either the x coordinate to be used to position the legend or a keyword accepted by legend() such as "topleft", "topright", "bottomleft", and "bottomright". |
|--------|--|
| У | numeric. The y coordinate to be used to position the legend. Can be NULL (default) if using keywords in \boldsymbol{x} . |
| legend | character. A character vector of the labels to appear in the legend. |
| img | A Picture or png array object, e.g., from using get_phylopic(). |
| name | character. A taxonomic name to be passed to get_uuid(). |
| uuid | character. A valid uuid for a PhyloPic silhouette (such as that returned by get_uuid() or pick_phylopic()). |
| ysize | [Deprecated] use the height argument instead. |
| height | numeric. Height of the legend silhouette(s). The width is determined by the aspect ratio of the original image. |
| color | character. Color of silhouette outline. If "original" or NA is specified, the original color of the silhouette outline will be used (usually the same as "transparent"). To remove the outline, you can set this to "transparent". |

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fill

character. Color of silhouette. If "original" is specified, the original color of the silhouette will be used (usually the same as "black"). If color is specified and fill is NA, color will be used as the fill color (for backwards compatibility). To remove the fill, you can set this to "transparent".

... Additional arguments passed to legend().

Details

This function can be used to add PhyloPic silhouettes as a legend to a base R plot. Arguments available in legend() can be used and passed via Note that not all arguments in legend() are compatible with add_phylopic_legend(). These include arguments for modifying lines (e.g. lty, lwd, seg.len), points (e.g. pch, pt.lwd), and shading (e.g. angle and density). This is due to add_phylopic_legend() using add_phylopic_base() to generate the legend symbols. However, arguments for adjusting the legend appearance such as text (e.g. cex), legend box (e.g. bg), and color (e.g. border) are compatible.

Examples

```
# Get UUIDs
uuids <- get_uuid(name = "Canis lupus", n = 2)
# Generate empty plot
plot(0:10, 0:10, type = "n", main = "Wolves")
# Add data points
add_phylopic_base(uuid = uuids,
    color = "black", fill = c("blue", "green"),
    x = c(2.5, 7.5), y = c(2.5, 7.5), height = 2)
# Add legend
add_phylopic_legend(uuid = uuids,
    height = 0.5, color = "black", fill = c("blue", "green"),
    x = "bottomright", legend = c("Wolf 1", "Wolf 2"),
    bg = "lightgrey")</pre>
```

browse_phylopic

Browse PhyloPic for a given taxonomic name or uuid

Description

This function provides a convenient way to browse PhyloPic for a given taxonomic name of uuid.

Usage

```
browse_phylopic(name = NULL, uuid = NULL)
```

Arguments

name character. A taxonomic name. Various taxonomic levels are supported (e.g. species, genus, family).

uuid character. A PhyloPic image uuid, as acquired by get_uuid().

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Details

This function returns a PhyloPic url for an input name or uuid and opens the user's default web browser at this url. If no name or uuid is supplied, the base url of PhyloPic images is returned and opened instead.

Value

A character vector of a valid PhyloPic url for the specified name or uuid. If no name or uuid is supplied, the base url of PhyloPic images is returned.

Examples

```
## Not run:
url <- browse_phylopic(name = "Acropora cervicornis")
## End(Not run)</pre>
```

flip_phylopic

Flip a PhyloPic silhouette along its horizontal and/or vertical axis

Description

The picture can be a Picture or png array object, e.g., from using get_phylopic().

Usage

```
flip_phylopic(img, horizontal = TRUE, vertical = FALSE)
```

Arguments

img A Picture or png array object, e.g., from using get_phylopic().

horizontal logical. Should the silhouette be flipped horizontally? vertical logical. Should the silhouette be flipped vertically?

Value

A Picture or png array object (matching the type of img)

See Also

```
Other transformations: recolor_phylopic(), rotate_phylopic()
```

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geom_phylopic

Geom for adding PhyloPic silhouettes to a plot

Description

This geom acts like ggplot2::geom_point(), except that the specified silhouettes are used as points. Silhouettes can be specified by their name, uuid, or image objects (img).

Usage

```
geom_phylopic(
  mapping = NULL,
  data = NULL,
  stat = "identity",
  position = "identity",
  ...,
  na.rm = FALSE,
  show.legend = FALSE,
  inherit.aes = TRUE,
  remove_background = TRUE,
  verbose = FALSE,
  filter = NULL
)
```

Arguments

mapping

Set of aesthetic mappings created by aes(). If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.

data

The data to be displayed in this layer. There are three options:

If NULL, the default, the data is inherited from the plot data as specified in the call to ggplot().

A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify() for which variables will be created.

A function will be called with a single argument, the plot data. The return value must be a data. frame, and will be used as the layer data. A function can be created from a formula (e.g. \sim head(.x, 10)).

stat

The statistical transformation to use on the data for this layer. When using a geom_*() function to construct a layer, the stat argument can be used the override the default coupling between geoms and stats. The stat argument accepts the following:

- A Stat ggproto subclass, for example StatCount.
- A string naming the stat. To give the stat as a string, strip the function name of the stat_ prefix. For example, to use stat_count(), give the stat as "count".

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 For more information and other ways to specify the stat, see the layer stat documentation.

position

A position adjustment to use on the data for this layer. This can be used in various ways, including to prevent overplotting and improving the display. The position argument accepts the following:

- The result of calling a position function, such as position_jitter(). This method allows for passing extra arguments to the position.
- A string naming the position adjustment. To give the position as a string, strip the function name of the position_ prefix. For example, to use position_jitter(), give the position as "jitter".
- For more information and other ways to specify the position, see the layer position documentation.

Other arguments passed on to layer()'s params argument. These arguments broadly fall into one of 4 categories below. Notably, further arguments to the position argument, or aesthetics that are required can *not* be passed through Unknown arguments that are not part of the 4 categories below are ignored.

- Static aesthetics that are not mapped to a scale, but are at a fixed value and apply to the layer as a whole. For example, colour = "red" or linewidth = 3. The geom's documentation has an **Aesthetics** section that lists the available options. The 'required' aesthetics cannot be passed on to the params. Please note that while passing unmapped aesthetics as vectors is technically possible, the order and required length is not guaranteed to be parallel to the input data.
- When constructing a layer using a stat_*() function, the ... argument can be used to pass on parameters to the geom part of the layer. An example of this is stat_density(geom = "area", outline.type = "both"). The geom's documentation lists which parameters it can accept.
- Inversely, when constructing a layer using a geom_*() function, the ... argument can be used to pass on parameters to the stat part of the layer.
 An example of this is geom_area(stat = "density", adjust = 0.5). The stat's documentation lists which parameters it can accept.
- The key_glyph argument of layer() may also be passed on through This can be one of the functions described as key glyphs, to change the display of the layer in the legend.

na.rm

If FALSE, the default, missing values are removed with a warning. If TRUE, missing values are silently removed.

show.legend

logical. Should this layer be included in the legends? FALSE, the default, never includes, NA includes if any aesthetics are mapped, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display.

inherit.aes

If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. borders().

remove_background

logical. Should any white background be removed from the silhouette(s)? See recolor_phylopic() for details.

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verbose logical. Should the attribution information for the used silhouette(s) be printed

to the console (see get_attribution())?

filter character. Filter by usage license if using the name aesthetic. Use "by" to limit

results to images which do not require attribution, "nc" for images which allows commercial usage, and "sa" for images without a ShareAlike clause. The user

can also combine these filters as a vector.

Details

One (and only one) of the img, name, or uuid aesthetics must be specified. The img aesthetic can be Picture objects or png array objects, e.g., from using get_phylopic(). Use the x and y aesthetics to place the silhouettes at specified positions on the plot. The height or width aesthetic specifies the height or width, respectively, of the silhouettes in the units of the y axis (only one is allowed). The aspect ratio of the silhouettes will always be maintained. The hjust and vjust aesthetics can be used to manage the justification of the silhouettes with respect to the x and y coordinates.

The color (default: NA), fill (default: "black"), and alpha (default: 1) aesthetics can be used to change the outline color, fill color, and transparency (outline and fill) of the silhouettes, respectively. If color is specified and fill is NA color will be used as the fill color (for backwards compatibility). If "original" is specified for the color aesthetic, the original color of the silhouette outline will be used (usually the same as "transparent"). If "original" is specified for the fill aesthetic, the original color of the silhouette body will be used (usually the same as "black"). To remove the fill or outline, you can set fill or color to "transparent", respectively.

The horizontal and vertical aesthetics can be used to flip the silhouettes. The angle aesthetic can be used to rotate the silhouettes. When specifying a horizontal and/or vertical flip **and** a rotation, the flip(s) will always occur first. If you would like to customize this behavior, you can flip and/or rotate the image within your own workflow using flip_phylopic() and rotate_phylopic().

Note that png array objects can only be rotated by multiples of 90 degrees. Also, outline colors do not currently work for png array objects.

Aesthetics

geom_phylopic understands the following aesthetics:

- x (required)
- y (required)
- img or uuid or name (one, and only one, required)
- height or width (optional, maximum of only one allowed)
- ysize [Deprecated] Deprecated in favor of height or width
- color or colour
- fill
- alpha
- horizontal
- vertical
- angle
- hjust

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 Learn more about setting these aesthetics in add_phylopic().

Examples

get_attribution

Get PhyloPic attribution data

Description

This function provides a convenient way to obtain attribution data for PhyloPic images via an image uuid returned by get_uuid().

Usage

```
get_attribution(uuid = NULL, img = NULL, text = FALSE, permalink = FALSE)
```

Arguments

| uuid | character. A vector of valid uuid(s) for PhyloPic silhouette(s) such as that returned by get_uuid() or pick_phylopic(). |
|-----------|---|
| img | A Picture or png array object from get_phylopic(). A list of these objects can also be supplied. If img is supplied, uuid is ignored. Defaults to NULL. |
| text | logical. Should attribution information be returned as a text paragraph? Defaults to FALSE. |
| permalink | logical. Should a permalink be created for this collection of uuid(s)? Defaults to FALSE. |

Details

This function returns image uuid specific attribution data, including: contributor name, contributor uuid, contributor contact, image uuid, license, and license abbreviation. If text is set to TRUE, a text paragraph with the contributor name, year of contribution, and license type is printed and image attribution data is returned invisibly (i.e. using invisible(). If permalink is set to TRUE, a permanent link (hosted by PhyloPic) will be generated. This link can be used to view and share details about the image silhouettes, including contributors and licenses.

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Value

A list of PhyloPic attribution data for an image uuid or a text output of relevant attribution information.

Examples

```
## Not run:
# Get valid uuid
uuid <- get_uuid(name = "Acropora cervicornis")
# Get attribution data for uuid
attri <- get_attribution(uuid = uuid)

# Get list of valid uuids
uuids <- get_uuid(name = "Scleractinia", n = 5)
# Get attribution data for uuids
get_attribution(uuid = uuids, text = TRUE)
# Get attribution data for uuids and create permalink
get_attribution(uuid = uuids, text = TRUE, permalink = TRUE)

## End(Not run)</pre>
```

get_phylopic

Retrieve an image for a given PhyloPic uuid

Description

This retrieves a PhyloPic silhouette as a vectorized or rasterized object.

Usage

```
get_phylopic(uuid = NULL, format = "vector", height = 512, preview = FALSE)
## S3 method for class 'Picture'
print(x, ...)
## S3 method for class 'phylopic'
print(x, ...)
```

| uuid | character. A PhyloPic image uuid. |
|---------|--|
| format | character. Format of the image. To return a vectorized image, use "vector". To return a rasterized image, use "raster" and specify a desired height. |
| height | numeric. If format is "raster", this is the desired height of the raster image in pixels. This is ignored if format is "vector". |
| preview | logical. If preview is TRUE, the returned image is plotted. Defaults to FALSE. |
| x | A Picture or png array object, e.g., from using get_phylopic(). |
| | Ignored |

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Details

The height argument is ignored if the format argument is not set to "raster". If format is "raster", the height argument specifies the height of the desired raster object. The width of this raster object will be determined by the original aspect ratio of the silhouette. If a pre-rendered raster exists with the desired dimensions, it will be downloaded from PhyloPic. If not, the vector image from PhyloPic will be used to render a raster object of the desired size.

Value

If format is "vector", a Picture object is returned. If format is "raster", a png array representing the rasterized image is returned. Either way, the uuid and download url are included as the "uuid" and "url" attributes, respectively.

Examples

```
## Not run:
# uuid
uuid <- "9fae30cd-fb59-4a81-a39c-e1826a35f612"

# Get data for an image
img_svg <- get_phylopic(uuid, format = "vector") # vector format
img_png <- get_phylopic(uuid, format = "raster") # raster format
## End(Not run)</pre>
```

get_uuid

Get a PhyloPic uuid

Description

This function provides a convenient way to obtain a valid uuid or image url for an input taxonomic name. As multiple silhouettes can exist for each species in PhyloPic, this function extracts the primary image.

Usage

```
get_uuid(name = NULL, img = NULL, n = 1, filter = NULL, url = FALSE)
```

| name | character. A taxonomic name. Various taxonomic levels are supported (e.g. species, genus, family). NULL can also be supplied which will skip the taxonomic filtering of the PhyloPic database. |
|------|--|
| img | A Picture or png array object from get_phylopic(). A list of these objects can also be supplied. If img is supplied, name and n are ignored. Defaults to NULL. |
| n | numeric. How many unids should be returned? Depending on the requested name, multiple silhouettes might exist. If n exceeds the number of available images, all available unids will be returned. This argument defaults to 1. |

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| filter character. Filter uuid(s) by usage license. Use "by" to limit results | to image |
|--|----------|
|--|----------|

uuids which do not require attribution, "nc" for image uuids which allow commercial usage, and "sa" for image uuids without a ShareAlike clause. The user

can also combine these filters as a vector.

url logical. If FALSE (default), only the uuid is returned. If TRUE, a valid PhyloPic

image url of the uuid is returned.

Details

This function returns uuid(s) or image url (svg) for an input name. If a specific image is desired, the user can make use of pick_phylopic to visually select the desired uuid/url.

Value

A character vector of a valid PhyloPic uuid or svg image url.

Examples

```
## Not run:
uuid <- get_uuid(name = "Acropora cervicornis")
uuid <- get_uuid(name = "Dinosauria", n = 5, url = TRUE)
## End(Not run)</pre>
```

phylopic_key_glyph

Use PhyloPics as legend key glyphs

Description

Specify existing images, taxonomic names, or PhyloPic uuids to use PhyloPic silhouettes as legend key glyphs in a ggplot plot.

Usage

```
phylopic_key_glyph(img = NULL, name = NULL, uuid = NULL)
```

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Details

One (and only one) of img, name, or uuid must be specified.

This argument may also be a vector/list of valid values if multiple silhouettes should be used as key glyphs. In this case, the specified silhouettes will be used as ordered as key glyphs one by one, with recycling as necessary.

Note that the sizes of the silhouettes in the legend are currently maximized based on the size of the key. This size can be modified using the legend.key.size argument in ggplot2::theme(). Therefore, the silhouettes will not reflect the size aesthetic, and this function should not be used for size legends.

Examples

pick_phylopic

Pick a PhyloPic image from available options

Description

This function provides a visually interactive way to pick an image and valid uuid for an input taxonomic name. As multiple silhouettes can exist for each organism in PhyloPic, this function is useful for choosing the right image/uuid for the user.

Usage

```
pick_phylopic(
  name = NULL,
  n = 5,
  uuid = NULL,
  view = 1,
  filter = NULL,
  auto = NULL
)
```

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Arguments

| name | character. A taxonomic name. Different taxonomic levels are supported (e.g. species, genus, family). |
|--------|---|
| n | numeric. How many unids should be viewed? Depending on the requested name, multiple silhouettes may exist. If n exceeds the number of available images, all available unids will be returned. Defaults to 5. Only relevant if name supplied. |
| uuid | character. A vector (or list) of valid PhyloPic silhouette uuids, such as that returned by get_uuid() or resolve_phylopic(). |
| view | numeric. Number of silhouettes that should be plotted at the same time. Defaults to 1. |
| filter | character. Filter uuid(s) by usage license. Use "by" to limit results to image uuids which do not require attribution, "nc" for image uuids which allow commercial usage, and "sa" for image uuids without a ShareAlike clause. The user can also combine these filters. Only relevant if name supplied. |
| auto | numeric. This argument allows the user to automate input into the menu choice. If the input value is 1, the first returned image will be selected. If the input value is 2, requested images will be automatically cycled through with the final image returned. If the input value is 3, a list of attribution information for each image is returned (this functionality is principally intended for testing). If NULL (default), the user must interactively respond to the called menu. |

Details

This function allows the user to visually select the desired image from a pool of silhouettes available for the input name.

Note that while the view argument can be any positive integer, weaker/older computers may have issues displaying very large numbers of images at the same time (i.e. view > 9). If no images are displayed in your plotting environment, try decreasing the value of view.

Value

A Picture object is returned. The uuid of the selected image is saved as the "uuid" attribute of the returned object and is also printed to console.

Examples

```
## Not run:
# Defaults pane layout
img <- pick_phylopic(name = "Canis lupus", n = 5)
# 3 x 3 pane layout
img <- pick_phylopic(name = "Scleractinia", n = 9, view = 9)
## End(Not run)</pre>
```

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plot.Picture

Preview a PhyloPic silhouette

Description

Preview a raster or vector representation of a PhyloPic silhouette. This will plot the silhouette on a new page in your default plotting device.

Usage

```
## S3 method for class 'Picture'
plot(x, ...)
## S3 method for class 'phylopic'
plot(x, ...)
```

Arguments

x A Picture or png array object, e.g., from using get_phylopic().... Other arguments passed on to grImport2::grid.picture() or grid::grid.raster().

recolor_phylopic

Recolor a PhyloPic image

Description

Function to recolor and change alpha levels of a PhyloPic image.

Usage

```
recolor_phylopic(
  img,
  alpha = 1,
  color = NULL,
  fill = NULL,
  remove_background = TRUE
)
```

Arguments

img A Picture or png array object, e.g., from using get_phylopic().

alpha numeric. A value between 0 and 1, specifying the opacity of the silhouette.

color character. Color to make the outline of the silhouette. If NULL, the outline color is not changed.

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fill character. Color to make the body of the silhouette. If NULL, the body color is not changed.

remove_background

logical. Should any white background be removed? Only useful if img is a Picture object. See details.

Details

Some PhyloPic silhouettes do not have a transparent background. Consequently, when color is used with vectorized versions of these images, the entire image—including the background—is recolored. Setting remove_background to TRUE (the default) will remove any white parts of the image (which should only be the background).

Value

A Picture or png array object (matching the type of img)

See Also

```
Other transformations: flip_phylopic(), rotate_phylopic()
```

resolve_phylopic

 $\label{thm:continuous} \textit{Use a taxonomic name from another database to get a PhyloPic image} \ \textit{UUID}$

Description

This function takes a supplied taxonomic name, queries it via the specified external API, resolves the API's returned taxonomic ID to the PhyloPic taxonomic node database, then retrieves PhyloPic image unids (or urls) for that node.

Usage

```
resolve_phylopic(
  name,
  api = "gbif.org",
  hierarchy = FALSE,
  max_ranks = 5,
  n = 1,
  filter = NULL,
  url = FALSE
)
```

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Arguments

| name | character. A full or partial taxonomic name to be queried via the specified api. |
|-----------|--|
| api | character. The API in which to query name. See Details for the available options. |
| hierarchy | logical. Whether the taxonomic hierarchy of name should be retrieved from the API and used to get several sets of PhyloPic image uuids (or urls). |
| max_ranks | numeric. The maximum number of taxonomic ranks that should be included if hierarchy is TRUE. |
| n | numeric. How many unids should be returned? Depending on the requested name, multiple silhouettes might exist. If n exceeds the number of available images, all available unids will be returned. This argument defaults to 1. |
| filter | character. Filter uuid(s) by usage license. Use "by" to limit results to image uuids which do not require attribution, "nc" for image uuids which allow commercial usage, and "sa" for image uuids without a ShareAlike clause. The user can also combine these filters as a vector. |
| url | logical. If FALSE (default), only the uuid is returned. If TRUE, a valid PhyloPic image url of the uuid is returned. |

Details

If hierarchy is FALSE, the specified name is queried via the specified api. The matched id is then resolved to the matching taxonomic node in the PhyloPic database. If hierarchy is TRUE, the full taxonomic hierarchy for name is retrieved from the specified api, those taxonomic names are subset to max_ranks ranks (starting from the specified name and ascending the hierarchy). Then each of those names is resolved to the matching taxonomic node in the PhyloPic database (where possible). In either case, get_uuid() is then used to retrieve n image UUID(s) for each taxonomic name.

Note that while the names of the returned list are the taxonomic names as reported by the specified api, the PhyloPic images that are returned are associated with whatever taxonomic node that taxonomic name resolves to in the PhyloPic database. This means that, if hierarchy is TRUE, the same images may be returned for multiple taxonomic names. Also, if a particular taxonomic name does not resolve to any node in the PhyloPic database, no images will be returned for that name.

The following APIs are available for querying (api):

- "eol.org": the Encyclopedia of Life (note: hierarchy = TRUE is not currently available for this API) ("eol" is also allowed)
- "gbif.org": the Global Biodiversity Information Facility ("gbif" is also allowed)
- "marinespecies.org": the World Registor of Marine Species ("worms" is also allowed)
- "opentreeoflife.org": the Open Tree of Life ("otol" is also allowed)
- "paleobiodb.org": the Paleobiology Database ("pbdb" is also allowed)

Value

A list where each value is n (or fewer) PhyloPic image uuids (or urls if url = TRUE) and each name is the taxonomic name as matched and reported by the specified api. If hierarchy is FALSE, the list has length 1. If hierarchy is TRUE, the list has maximum length max_ranks.

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Examples

rotate_phylopic

Rotate a PhyloPic silhouette

Description

The picture can be a Picture or png array object, e.g., from using get_phylopic(). Note that png array objects can only be rotated by multiples of 90 degrees.

Usage

```
rotate_phylopic(img, angle = 90)
```

Arguments

img A Picture or png array object, e.g., from using get_phylopic().
angle numeric. The number of degrees to rotate the silhouette clockwise.

Value

A Picture or png array object (matching the type of img)

See Also

Other transformations: flip_phylopic(), recolor_phylopic()

save_phylopic

Save a PhyloPic image

Description

save_phylopic is a convenient function for saving a PhyloPic silhouette fetched via get_phylopic. Available formats for saving are those available in the grDevices package (i.e. pdf, png, svg, tiff, jpeg, and bmp).

Usage

```
save_phylopic(img = NULL, path = NULL, bg = "transparent", ...)
```

Arguments

| img | picture. A Picture object. For PhyloPic silhouettes, this is generated by get_phylopic and pick_phylopic. |
|------|--|
| path | character. The output file path. The name of the file and file extension should be included (e.g. "./phylopic.png"). If this argument is NULL (default), the file is automatically saved in the current working directory as a png file. |
| bg | character. The desired background color. By default this argument is set to "transparent". Note: .jpeg files are always exported with a background colour. |
| | Additional arguments available for the called grDevices function. The called function is determined by the file extension used in path (e.g. png). |

Details

This function can be used to save a PhyloPic image obtained via the get_phylopic or any object of class Picture from the grImport2 package. Any file format available in the grDevices package can be used (i.e. pdf, png, svg, tiff, jpeg, and bmp).

Value

An image is saved. Nothing is returned to the R interface.

Examples

```
scale_height_continuous
```

Scales for phylopic height or width

Description

scale_height_continuous() scales the height of silhouettes whereas scale_width_continuous() scales the height of silhouettes.

Usage

```
scale_height_continuous(
  name = waiver(),
  breaks = waiver(),
  labels = waiver(),
  limits = NULL,
  range = c(1, 6),
  transform = "identity",
  trans = lifecycle::deprecated(),
  guide = "none"
scale_width_continuous(
  name = waiver(),
  breaks = waiver(),
  labels = waiver(),
  limits = NULL,
  range = c(1, 6),
  transform = "identity",
  trans = lifecycle::deprecated(),
  guide = "none"
)
```

Arguments

name

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

breaks One of:

- · NULL for no breaks
- waiver() for the default breaks computed by the transformation object
- A numeric vector of positions
- A function that takes the limits as input and returns breaks as output (e.g., a function returned by scales::extended_breaks()). Note that for position scales, limits are provided after scale expansion. Also accepts rlang lambda function notation.

labels One of:

- NULL for no labels
- waiver() for the default labels computed by the transformation object
- A character vector giving labels (must be same length as breaks)
- An expression vector (must be the same length as breaks). See ?plotmath for details.
- A function that takes the breaks as input and returns labels as output. Also accepts rlang lambda function notation.

limits One of:

- NULL to use the default scale range
- A numeric vector of length two providing limits of the scale. Use NA to refer to the existing minimum or maximum
- A function that accepts the existing (automatic) limits and returns new limits. Also accepts rlang lambda function notation. Note that setting limits on positional scales will **remove** data outside of the limits. If the purpose is to zoom, use the limit argument in the coordinate system (see coord_cartesian()).

range

a numeric vector of length 2 that specifies the minimum and maximum size of the plotting symbol after transformation.

transform

For continuous scales, the name of a transformation object or the object itself. Built-in transformations include "asn", "atanh", "boxcox", "date", "exp", "hms", "identity", "log", "log10", "log1p", "log2", "logit", "modulus", "probability", "probit", "pseudo_log", "reciprocal", "reverse", "sqrt" and "time".

A transformation object bundles together a transform, its inverse, and methods for generating breaks and labels. Transformation objects are defined in the scales package, and are called transform_<name>. If transformations require arguments, you can call them from the scales package, e.g. scales::transform_boxcox(p = 2). You can create your own transformation with scales::new_transform().

trans

[Deprecated] Deprecated in favour of transform.

guide

A function used to create a guide or its name. See guides() for more information.

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