

Package ‘ggquickeda’

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Title Quickly Explore Your Data Using 'ggplot2' and 'table1' Summary Tables

Version 0.3.2

Description Quickly and easily perform exploratory data analysis by uploading your data as a 'csv' file. Start generating insights using 'ggplot2' plots and 'table1' tables with descriptive stats, all using an easy-to-use point and click 'Shiny' interface.

URL <https://github.com/smouksassi/ggquickeda>,
<https://smouksassi.github.io/ggquickeda/>

BugReports <https://github.com/smouksassi/ggquickeda/issues>

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geom_km	<i>Add a Kaplan-Meier survival curve</i>
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Description

Add a Kaplan-Meier survival curve

Usage

```
geom_km(
  mapping = NULL,
  data = NULL,
  stat = "km",
  position = "identity",
  show.legend = NA,
  inherit.aes = TRUE,
  na.rm = TRUE,
  ...
)
```

Arguments

mapping	Set of aesthetic mappings created by aes() . If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply <code>mapping</code> if there is no plot mapping.
data	The data to be displayed in this layer. There are three options: If <code>NULL</code> , the default, the data is inherited from the plot data as specified in the call to ggplot() . A <code>data.frame</code> , 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 <code>data.frame</code> , and will be used as the layer data. A function can be created from a formula (e.g. <code>~ head(.x, 10)</code>).
stat	The statistical transformation to use on the data for this layer. When using a <code>geom_*</code> function to construct a layer, the <code>stat</code> argument can be used to override the default coupling between geoms and stats. The <code>stat</code> argument accepts the following: <ul style="list-style-type: none">• A Stat ggproto subclass, for example <code>StatCount</code>.• A string naming the stat. To give the stat as a string, strip the function name of the <code>stat_</code> prefix. For example, to use <code>stat_count()</code>, give the stat as "count".• 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 <code>position</code> argument accepts the following: <ul style="list-style-type: none">• The result of calling a position function, such as <code>position_jitter()</code>. 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 <code>position_</code> prefix. For example, to use <code>position_jitter()</code>, give the position as "jitter".• For more information and other ways to specify the position, see the layer position documentation.
show.legend	logical. Should this layer be included in the legends? <code>NA</code> , the default, includes if any aesthetics are mapped. <code>FALSE</code> never includes, and <code>TRUE</code> always includes. It can also be a named logical vector to finely select the aesthetics to display. To include legend keys for all levels, even when no data exists, use <code>TRUE</code> . If <code>NA</code> , all levels are shown in legend, but unobserved levels are omitted.
inherit.aes	If <code>FALSE</code> , 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. annotation_borders() .
na.rm	If <code>FALSE</code> , the default, missing values are removed with a warning. If <code>TRUE</code> , missing values are silently removed.

...

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.

Aesthetics

`geom_km` understands the following aesthetics (required aesthetics are in bold):

- `x` The survival/censoring times. This is automatically mapped by `stat_km()`
- `y` The survival probability estimates. This is automatically mapped by `stat_km()` smallest level in sort order is assumed to be 0, with a warning.
- `alpha`
- `color`
- `linetype`
- `size`

See Also

The default stat for this geom is `stat_km()` see that documentation for more options to control the underlying statistical transformation.

Examples

```
library(ggplot2)
set.seed(123)
sex <- rbinom(250, 1, .5)
df <- data.frame(time = exp(rnorm(250, mean = sex)), status = rbinom(250, 1, .75), sex = sex)
ggplot(df, aes(time = time, status = status, color = factor(sex))) + geom_km()
```

`geom_kmband`*Add confidence bands to a Kaplan-Meier survival curve*

Description

Add confidence bands to a Kaplan-Meier survival curve

Usage

```
geom_kmband(  
  mapping = NULL,  
  data = NULL,  
  stat = "kmband",  
  position = "identity",  
  show.legend = NA,  
  inherit.aes = TRUE,  
  na.rm = TRUE,  
  ...  
)
```

Arguments

<code>mapping</code>	Set of aesthetic mappings created by aes() . If specified and <code>inherit.aes</code> = <code>TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply <code>mapping</code> if there is no plot mapping.
<code>data</code>	The data to be displayed in this layer. There are three options: If <code>NULL</code> , the default, the data is inherited from the plot data as specified in the call to ggplot() . A <code>data.frame</code> , 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 <code>data.frame</code> , and will be used as the layer data. A function can be created from a <code>formula</code> (e.g. <code>~ head(.x, 10)</code>).
<code>stat</code>	The statistical transformation to use on the data for this layer. When using a <code>geom_*</code> function to construct a layer, the <code>stat</code> argument can be used to override the default coupling between geoms and stats. The <code>stat</code> argument accepts the following: <ul style="list-style-type: none">• A Stat ggproto subclass, for example <code>StatCount</code>.• A string naming the stat. To give the stat as a string, strip the function name of the <code>stat_</code> prefix. For example, to use <code>stat_count()</code>, give the stat as <code>"count"</code>.• 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: <ul style="list-style-type: none"> The result of calling a position function, such as <code>position_jitter()</code>. 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 <code>position_</code> prefix. For example, to use <code>position_jitter()</code>, give the position as "jitter". For more information and other ways to specify the position, see the layer position documentation.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display. To include legend keys for all levels, even when no data exists, use TRUE. If NA, all levels are shown in legend, but unobserved levels are omitted.
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. annotation_borders() .
na.rm	If FALSE, the default, missing values are removed with a warning. If TRUE, missing values are silently removed.
...	Other arguments passed on to <code>layer()</code> 's <code>params</code> argument. These arguments broadly fall into one of 4 categories below. Notably, further arguments to the <code>position</code> argument, or aesthetics that are required can <i>not</i> be passed through Unknown arguments that are not part of the 4 categories below are ignored. <ul style="list-style-type: none"> 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, <code>colour = "red"</code> or <code>linewidth = 3</code>. The geom's documentation has an Aesthetics section that lists the available options. The 'required' aesthetics cannot be passed on to the <code>params</code>. 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 <code>stat_*</code>() function, the ... argument can be used to pass on parameters to the geom part of the layer. An example of this is <code>stat_density(geom = "area", outline.type = "both")</code>. The geom's documentation lists which parameters it can accept. Inversely, when constructing a layer using a <code>geom_*</code>() function, the ... argument can be used to pass on parameters to the stat part of the layer. An example of this is <code>geom_area(stat = "density", adjust = 0.5)</code>. The stat's documentation lists which parameters it can accept. The <code>key_glyph</code> argument of <code>layer()</code> 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.

Aesthetics

`geom_kmband` understands the following aesthetics (required aesthetics are in bold):

- `x` The survival/censoring times. This is automatically mapped by `stat_kmband()`
- `y` The survival probability estimates. This is automatically mapped by `stat_kmband()` smallest level in sort order is assumed to be 0, with a warning
- `alpha`
- `color`
- `linetype`
- `linewidth`

See Also

The default stat for this geom is `stat_kmband()`. See that documentation for more options to control the underlying statistical transformation.

Examples

```
library(ggplot2)
sex <- rbinom(250, 1, .5)
df <- data.frame(time = exp(rnorm(250, mean = sex)), status = rbinom(250, 1, .75), sex = sex)
ggplot(df, aes(time = time, status = status, color = factor(sex), fill = factor(sex))) +
  geom_km() + geom_kmband()
```

geom_kmticks

Add tick marks to a Kaplan-Meier survival curve

Description

Adds tickmarks at the times when there are censored observations but no events

Usage

```
geom_kmticks(
  mapping = NULL,
  data = NULL,
  stat = "kmticks",
  position = "identity",
  show.legend = NA,
  inherit.aes = TRUE,
  na.rm = TRUE,
  ...
)
```

Arguments

<code>mapping</code>	Set of aesthetic mappings created by <code>aes()</code> . If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply <code>mapping</code> if there is no plot mapping.
<code>data</code>	The data to be displayed in this layer. There are three options: If <code>NULL</code> , the default, the data is inherited from the plot data as specified in the call to <code>ggplot()</code> . A <code>data.frame</code> , or other object, will override the plot data. All objects will be fortified to produce a data frame. See <code>fortify()</code> for which variables will be created. A function will be called with a single argument, the plot data. The return value must be a <code>data.frame</code> , and will be used as the layer data. A function can be created from a formula (e.g. <code>~ head(.x, 10)</code>).
<code>stat</code>	The statistical transformation to use on the data for this layer. When using a <code>geom_*</code> function to construct a layer, the <code>stat</code> argument can be used to override the default coupling between geoms and stats. The <code>stat</code> argument accepts the following: <ul style="list-style-type: none"> • A Stat ggproto subclass, for example <code>StatCount</code>. • A string naming the stat. To give the stat as a string, strip the function name of the <code>stat_</code> prefix. For example, to use <code>stat_count()</code>, give the stat as "count". • For more information and other ways to specify the stat, see the layer stat documentation.
<code>position</code>	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 <code>position</code> argument accepts the following: <ul style="list-style-type: none"> • The result of calling a position function, such as <code>position_jitter()</code>. 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 <code>position_</code> prefix. For example, to use <code>position_jitter()</code>, give the position as "jitter". • For more information and other ways to specify the position, see the layer position documentation.
<code>show.legend</code>	logical. Should this layer be included in the legends? <code>NA</code> , the default, includes if any aesthetics are mapped. <code>FALSE</code> never includes, and <code>TRUE</code> always includes. It can also be a named logical vector to finely select the aesthetics to display. To include legend keys for all levels, even when no data exists, use <code>TRUE</code> . If <code>NA</code> , all levels are shown in legend, but unobserved levels are omitted.
<code>inherit.aes</code>	If <code>FALSE</code> , 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. <code>annotation_borders()</code> .
<code>na.rm</code>	If <code>FALSE</code> , the default, missing values are removed with a warning. If <code>TRUE</code> , missing values are silently removed.

...

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.

Aesthetics

`geom_kmticks` understands the following aesthetics (required aesthetics are in bold):

- `x` The survival/censoring times. This is automatically mapped by `stat_kmticks()`
- `y` The survival probability estimates. This is automatically mapped by `stat_kmticks()` smallest level in sort order is assumed to be 0, with a warning
- `alpha`
- `color`
- `linetype`
- `size`

See Also

The default stat for this geom is `stat_kmticks` see that documentation for more options to control the underlying statistical transformation.

Examples

```
library(ggplot2)
sex <- rbinom(250, 1, .5)
df <- data.frame(time = exp(rnorm(250, mean = sex)), status = rbinom(250, 1, .75), sex = sex)
ggplot(df, aes(time = time, status = status, color = factor(sex), group = factor(sex))) +
  geom_km() + geom_kmticks(col="black")
```

ggcontinuousexpdist *Create a continuous fit vs exposure(s) plot*

Description

Produces a logistic fit plot with a facetable exposures/quantiles/distributions in ggplot2

Usage

```
ggcontinuousexpdist(
  data = effICGI,
  response = "response",
  endpoint = "Endpoint",
  model_type = c("linear", "loess", "none"),
  DOSE = "DOSE",
  color_fill = "DOSE",
  fit_by_color_fill = FALSE,
  exposure_metrics = c("AUC", "CMAX"),
  exposure_metric_split = c("median", "tertile", "quartile", "none"),
  exposure_metric_soc_value = -99,
  exposure_metric_plac_value = 0,
  exposure_distribution = c("distributions", "lineranges", "boxplots", "none"),
  exposure_distribution_percent = TRUE,
  exposure_distribution_percent_text_size = 5,
  dose_plac_value = "Placebo",
  xlab = "Exposure Values",
  ylab = "Response",
  points_alpha = 0.2,
  points_show = TRUE,
  mean_obs_byexptile = TRUE,
  mean_obs_byexptile_plac = TRUE,
  mean_obs_byexptile_group = "none",
  mean_text_size = 5,
  mean_obs_bydose = TRUE,
  mean_obs_bydose_plac = FALSE,
  N_text_show = TRUE,
  N_text_size = 5,
  N_text_ypos = c("with means", "top"),
  N_text_sep = "\n",
  binlimits_show = TRUE,
  binlimits_text_size = 5,
  binlimits_ypos = -Inf,
  binlimits_color = "#B3B3B380",
  dist_position_scaler = 0.2,
  dist_offset = 0,
  dist_scale = 0.9,
  lineranges_ypos = -1,
```

```

    lineranges_dodge = 1,
    lineranges_doselabel = FALSE,
    proj_bydose = TRUE,
    yproj = TRUE,
    yproj_xpos = 0,
    yproj_dodge = 0.2,
    yaxis_position = c("left", "right"),
    facet_formula = NULL,
    theme_certara = TRUE,
    return_list = FALSE
)

```

Arguments

<code>data</code>	Data to use with multiple endpoints stacked into response (values), Endpoint(endpoint name)
<code>response</code>	name of the column holding the response values
<code>endpoint</code>	name of the column holding the name/key of the endpoint default to Endpoint
<code>model_type</code>	type of the trend fit one of "linear", "loess", "none"
<code>DOSE</code>	name of the column holding the DOSE values default to DOSE
<code>color_fill</code>	name of the column to be used for color/fill default to DOSE column
<code>fit_by_color_fill</code>	fit split by color? default FALSE
<code>exposure_metrics</code>	<p>name(s) of the column(s) to be stacked into expname exptile and split into exposure_metric_split</p>
<code>exposure_metric_split</code>	one of "median", "tertile", "quartile", "none"
<code>exposure_metric_soc_value</code>	special exposure code for standard of care default -99
<code>exposure_metric_plac_value</code>	special exposure code for placebo default 0
<code>exposure_distribution</code>	one of distributions, lineranges, boxplots or none
<code>exposure_distribution_percent</code>	show percent of distribution between binlimits TRUE/FALSE
<code>exposure_distribution_percent_text_size</code>	distribution percentages text size default to 5
<code>dose_plac_value</code>	string identifying placebo in DOSE column
<code>xlab</code>	text to be used as x axis label
<code>ylab</code>	text to be used as y axis label
<code>points_alpha</code>	alpha transparency for points
<code>points_show</code>	show the observations TRUE/FALSE

```

mean_obs_byexptile
    observed mean by exptile TRUE/FALSE
mean_obs_byexptile_plac
    observed mean by exptile placebo TRUE/FALSE
mean_obs_byexptile_group
    additional grouping for exptile means default none
mean_text_size  mean text size default to 5
mean_obs_bydose
    observed mean by dose TRUE/FALSE
mean_obs_bydose_plac
    observed mean by placebo dose TRUE/FALSE
N_text_show     show the N by exptile TRUE/FALSE
N_text_size     N by exposure text size default to 5
N_text_ypos     y position for N two text elements the first for by exptile and the second for by dose/color options include with means top bottom
N_text_sep      character string to separate N from mean default \n
binlimits_show  show the binlimits vertical lines TRUE/FALSE
binlimits_text_size
    binlimits text size default to 5
binlimits_ypos  binlimits y position default to -Inf
binlimits_color
    binlimits text color default to alpha("gray70",0.5)
dist_position_scaler
    space occupied by the distribution default to 0.2
dist_offset      offset where the distribution position starts default to 0
dist_scale       scaling parameter for ggridges default to 0.9
lineranges_ypos
    where to put the lineranges -1
lineranges_dodge
    lineranges vertical dodge value 1
lineranges_doselabel
    TRUE/FALSE
proj_bydose     project the predictions on logistic curve TRUE/FALSE
yproj           project the predictions on y axis TRUE/FALSE
yproj_xpos      y projection x position 0
yproj_dodge     y projection dodge value 0.2
yaxis_position  where to put y axis "left" or "right"
facet_formula   facet formula to be use otherwise endpoint ~ expname
theme_certara  apply certara colors and format for strips and default colour/fill
return_list     What to return if True a list of the datasets and plot is returned instead of only the plot

```

Examples

```

# Example 1
library(ggplot2)
library(patchwork)
effICGI <- logistic_data |>
dplyr::filter(!is.na(ICGI7))|>
dplyr::filter(!is.na(AUC))
effICGI$DOSE <- factor(effICGI$DOSE,
levels=c("0", "600", "1200","1800","2400"),
labels=c("Placebo", "600 mg", "1200 mg","1800 mg","2400 mg"))
effICGI$STUDY <- factor(effICGI$STUDY)
effICGI <- tidyr::gather(effICGI,Endpoint,response,ICGI7,BRLS)
a <- ggcontinuousexpdist(data = effICGI |> dplyr::filter(Endpoint =="ICGI7"),
  response = "response",
  endpoint = "Endpoint",
  exposure_metrics = c("AUC"),
  exposure_metric_split = c("quartile"),
  exposure_metric_soc_value = -99,
  exposure_metric_plac_value = 0,
  dist_position_scaler = 1, dist_offset = -1 ,
  yproj_xpos = -20 ,
  yproj_dodge = 20 ,
  exposure_distribution ="distributions",
  mean_obs_bydose_plac = TRUE,
  mean_obs_byexptile_plac=FALSE,
  return_list = FALSE
)

b <- ggcontinuousexpdist(data = effICGI |> dplyr::filter(Endpoint =="BRLS"),
  response = "response",
  endpoint = "Endpoint",
  exposure_metrics = c("AUC"),
  exposure_metric_split = c("quartile"),
  exposure_metric_soc_value = -99,
  exposure_metric_plac_value = 0,
  dist_position_scaler = 4.2, dist_offset = 5 ,
  yproj_xpos = -20 ,
  yproj_dodge = 20 ,
  exposure_distribution ="distributions",
  mean_obs_bydose_plac = TRUE,
  mean_obs_byexptile_plac=FALSE,
  return_list = FALSE)
(a / b) +
plot_layout(guides = "collect") &
theme(legend.position = "top")

#Example 2 loess fit
effICGI$SEX <- as.factor(effICGI$SEX)
ggcontinuousexpdist(data = effICGI |>
  dplyr::filter(Endpoint =="ICGI7"),
  model_type = "loess",
  response = "response",

```

```

endpoint = "Endpoint",
color_fill = "SEX",
exposure_metrics = c("AUC"),
exposure_metric_split = c("quartile"),
exposure_metric_soc_value = -99,
exposure_metric_plac_value = 0,
dist_position_scaler = 1, dist_offset = -1 ,
yproj_xpos = -20 ,
yproj_dodge = 20 ,
exposure_distribution ="lineranges",
lineranges_ypos = -0.2,
lineranges_dodge = 0.2,
lineranges_doselabel = TRUE)

## Not run:
#Example 3
library(ggplot2)
library(patchwork)
effICGI <- logistic_data |>
dplyr::filter(!is.na(ICGI7))|>
dplyr::filter(!is.na(AUC))
effICGI$DOSE <- factor(effICGI$DOSE,
levels=c("0", "600", "1200", "1800", "2400"),
labels=c("Placebo", "600 mg", "1200 mg", "1800 mg", "2400 mg"))
effICGI$STUDY <- factor(effICGI$STUDY)
effICGI <- tidyverse::gather(effICGI, Endpoint, response, PRLS, BRLS)
effICGI$Endpoint2 <- effICGI$Endpoint
ggcontinuousexpdist(data = effICGI ,
model_type = "loess",
response = "response",
endpoint = "Endpoint",
color_fill = "Endpoint2",
fit_by_color_fill = TRUE,
exposure_metrics = c("AUC"),
exposure_metric_split = c("quartile"),
exposure_metric_soc_value = -99,
exposure_metric_plac_value = 0,
yproj = FALSE, points_show = FALSE,
exposure_distribution ="none",
N_text_sep = " | N = ",
N_text_ypos = c("bottom", "with means"),
binlimits_color = "red",
binlimits_ypos = 10,
facet_formula = ~expname)

## End(Not run)

```

Description

Produces a km plot with a facettable risk table in ggplot2

Usage

```
ggkmrisktable(  
  data = lung_long,  
  time = "time",  
  status = "DV",  
  endpoint = "Endpoint",  
  groupvar1 = "Endpoint",  
  groupvar2 = "expname",  
  groupvar3 = "none",  
  exposure_metrics = c("age", "ph.karno"),  
  exposure_metric_split = c("median", "tertile", "quartile", "none"),  
  exposure_metric_soc_value = -99,  
  exposure_metric_plac_value = 0,  
  exposure_metric_soc_name = "SOC",  
  exposure_metric_plac_name = "Placebo",  
  show_exptile_values = FALSE,  
  show_exptile_values_pos = c("left", "right"),  
  show_exptile_values_textsize = 5,  
  show_exptile_values_order = c("default", "reverse"),  
  color_fill = "exptile",  
  linetype = "exptile",  
  xlab = "Time of follow_up",  
  ylab = "Overall survival probability",  
  nrisk_table_plot = TRUE,  
  nrisk_table_variables = c("n.risk", "pct.risk", "n.event", "cum.n.event", "n.censor"),  
  nrisk_table_breaktimeby = NULL,  
  nrisk_table_textsize = 4,  
  nrisk_position_scaler = 0.2,  
  nrisk_position_dodge = 0.2,  
  nrisk_offset = 0,  
  nrisk_filterout0 = FALSE,  
  km_logrank_pvalue = FALSE,  
  km_logrank_pvalue_pos = c("left", "right"),  
  km_logrank_pvalue_textsize = 5,  
  km_logrank_pvalue_cutoff = 0.001,  
  km_logrank_pvalue_digits = 3,  
  km_trans = c("identity", "event", "cumhaz", "cloglog"),  
  km_linewidth = 1,  
  km_ticks = TRUE,  
  km_band = TRUE,  
  km_conf_int = 0.95,  
  km_conf_type = c("log", "plain", "log", "log-log", "logit", "none"),  
  km_conf_lower = c("usual", "peto", "modified"),  
  km_median = c("none", "median", "medianci", "table"),  
  km_median_table_pos = c("left", "right"),  
  km_median_table_order = c("default", "reverse"),  
  km_yaxis_position = c("left", "right"),  
  facet_formula = NULL,
```

```

    facet_ncol = NULL,
    facet_strip_position = c("top", "top", "top", "top"),
    theme_certara = TRUE,
    return_list = FALSE
)

```

Arguments

data	Data to use with multiple endpoints stacked into time, status, endpoint name
time	name of the column holding the time to event information default to time
status	name of the column holding the event information default to DV
endpoint	name of the column holding the name/key of the endpoint default to Endpoint
groupvar1	name of the column to group by, default Endpoint
groupvar2	name of the column to group by in addition to groupvar1, default expname
groupvar3	name of the column to group by in addition to groupvar1 and groupvar2, default "none"
exposure_metrics	name(s) of the column(s) to be stacked into expname exptile and split into exposure_metric_split
exposure_metric_split	one of "median", "tertile", "quartile", "none"
exposure_metric_soc_value	special exposure code for standard of care default -99
exposure_metric_plac_value	special exposure code for placebo default 0
exposure_metric_soc_name	soc name default to "soc"
exposure_metric_plac_name	placebo name default to "placebo"
show_exptile_values	FALSE
show_exptile_values_pos	"left" or "right"
show_exptile_values_textsize	default to 5
show_exptile_values_order	the order of the entries "default" or "reverse"
color_fill	name of the column to be used for color/fill default to exptile
linetype	name of the column to be used for linetype default to exptile
xlab	text to be used as x axis label
ylab	text to be used as y axis label
nrisk_table_plot	TRUE

```

nrisk_table_variables
  one or more from: "n.risk", "pct.risk", "n.event", "cum.n.event", "n.censor"
nrisk_table_breaktimeby
  NULL
nrisk_table_textsize
  4
nrisk_position_scaler
  0.2
nrisk_position_dodge
  0.2, negative values will reverse the order
nrisk_offset  0
nrisk_filterout0
  FALSE
km_logrank_pvalue
  FALSE
km_logrank_pvalue_pos
  "left" or "right"
km_logrank_pvalue_textsize
  pvalue text size default to 5
km_logrank_pvalue_cutoff
  pvalue below which to print as p < cutoff
km_logrank_pvalue_digits
  pvalue ndigits for round function
km_trans      one of "identity", "event", "cumhaz", "cloglog"
km_linewidth   linewidth for the km curves default to 1
km_ticks       TRUE
km_band        TRUE
km_conf_int    0.95
km_conf_type   default one of "log", "plain", "log-log", "logit", "none"
km_conf_lower  one of "usual", "peto", "modified"
km_median      add median survival information one of "none", "median", "medianci", "table"
km_median_table_pos
  when table is chosen where to put it "left" or "right"
km_median_table_order
  when table is chosen the order of the entries "default" or "reverse"
km_yaxis_position
  where to put y axis on "left" or "right"
facet_formula facet formula to be used otherwise ~ groupvar1 + groupvar2 + groupvar3
facet_ncol    NULL if not specified the automatic waiver will be used
facet_strip_position
  position in sequence for the variable used in faceting default to c("top", "top", "top", "top")
theme_certara apply certara colors and format for strips and default colour/fill
return_list    What to return if True a list of the datasets and plot is returned instead of only
               the plot

```

Examples

```

library(tidyr)
# Example 1
lung_long <- survival::lung |>
  dplyr::mutate(status = ifelse(status==1,0,1)) |>
  tidyr::gather(Endpoint,DV,status) |>
  dplyr::filter(!is.na(ph.karno))|>
  dplyr::filter(!is.na(pat.karno))|>
  dplyr::filter(!is.na(ph.ecog))
lung_long$ph.ecog <- ifelse(lung_long$ph.ecog>1,2,lung_long$ph.ecog)
lung_long$ph.ecog <- as.factor(lung_long$ph.ecog )
lung_long$ph.ecog <- as.factor(lung_long$ph.ecog )
lung_long$facetdum <- "(all)"

ggkmrisktable(data = lung_long, time= "time", status ="DV",
               exposure_metrics =c("age","ph.karno"),
               exposure_metric_split = "tertile",
               color_fill = "exptile",
               linetype = "expname",
               groupvar1 = "Endpoint",
               groupvar2 = "exptile",
               xlab = "Time of follow_up",
               ylab ="Overall survival probability",
               nrisk_table_variables = c("n.risk","n.event"),
               km_median = "medianci",
               km_band = FALSE,
               nrisk_table_breaktimeby = 200,
               facet_ncol = 3)

#Example 2
ggkmrisktable(data = lung_long, time= "time", status ="DV",
               exposure_metrics =c("age","ph.karno"),
               exposure_metric_split = "quartile",
               color_fill = "exptile",
               linetype = "none",
               groupvar1 = "Endpoint",
               groupvar2 = "exptile",
               xlab = "Time of follow_up",
               ylab ="Overall survival probability",
               nrisk_table_variables = c("cum.n.event","pct.risk","n.censor"),
               km_median = "medianci",
               km_band = TRUE,
               km_trans = "event",
               show_exptile_values = TRUE,
               show_exptile_values_pos = "right",
               nrisk_table_breaktimeby = 200,
               facet_ncol = 3,
               facet_formula = ~expname)

## Not run:
#Example 3
ggkmrisktable(data = lung_long, time = "time", status = "DV",
               exposure_metrics =c("ph.karno","pat.karno"),
               exposure_metric_split = "median",

```

```

color_fill = "exptile",
linetype = "exptile",
groupvar1 = "Endpoint",
groupvar2 = "expname",
xlab = "Time of follow_up",
ylab ="Overall survival probability",
nrisk_table_variables = c("n.event"),
km_trans = "event",
km_median = "table",
km_median_table_pos = "right",
km_logrank_pvalue = TRUE,
km_logrank_pvalue_cutoff = 0.0001,
km_logrank_pvalue_digits = 3,
km_band = TRUE,
nrisk_table_breaktimeby = 200,
facet_ncol = 3,
facet_formula = ~expname)

#Example 4
ggkmrisktable(data=lung_long,
               exposure_metrics = c("ph.karno", "age"),
               exposure_metric_split = "median",
               time = "time",
               status ="DV",
               color_fill = "ph.ecog",
               linetype = "ph.ecog",
               groupvar1 = "Endpoint",
               groupvar2 = "expname",
               groupvar3 = "exptile",
               nrisk_filterout0 = FALSE,
               nrisk_table_breaktimeby = 200,
               km_logrank_pvalue = TRUE,
               km_median = "table",
               km_median_table_pos = "left",
               facet_formula = ~expname+exptile)

#Example 5
ggkmrisktable(data=lung_long,
               exposure_metrics = c("ph.karno", "age"),
               exposure_metric_split = "none",
               color_fill = "none",
               linetype = "none",
               nrisk_table_variables = c("n.risk", "pct.risk", "n.event", "cum.n.event", "n.censor"),
               km_median = "table",
               nrisk_position_scaler = 0.1
               )

## End(Not run)

```

Description

Produces a logistic fit plot with a facetable exposures/quantiles/distributions in ggplot2

Usage

```
gglogisticexpdist(
  data = effICGI,
  response = "response",
  endpoint = "Endpoint",
  model_type = c("logistic"),
  DOSE = "DOSE",
  color_fill = "DOSE",
  logistic_by_color_fill = FALSE,
  exposure_metrics = c("AUC", "CMAX"),
  exposure_metric_split = c("median", "tertile", "quartile", "none"),
  exposure_metric_soc_value = -99,
  exposure_metric_plac_value = 0,
  exposure_distribution = c("distributions", "lineranges", "boxplots", "none"),
  exposure_distribution_percent = TRUE,
  exposure_distribution_percent_text_size = 5,
  dose_plac_value = "Placebo",
  xlab = "Exposure Values",
  ylab = "Probability of Response",
  points_alpha = 0.2,
  points_show = TRUE,
  prob_obs_byexptile = TRUE,
  prob_obs_byexptile_plac = TRUE,
  prob_obs_byexptile_group = "none",
  prob_text_size = 5,
  prob_obs_bydose = TRUE,
  prob_obs_bydose_plac = FALSE,
  Nresp_Ntot_show = TRUE,
  Nresp_Ntot_size = 5,
  Nresp_Ntot_ypos = c("with percentages", "top"),
  Nresp_Ntot_sep = "/",
  binlimits_show = TRUE,
  binlimits_text_size = 5,
  binlimits_ypos = 0,
  binlimits_color = "#B3B3B380",
  dist_position_scaler = 0.2,
  dist_offset = 0,
  dist_scale = 0.9,
  lineranges_ypos = 0.2,
  lineranges_dodge = 0.15,
  lineranges_doselabel = FALSE,
  proj_bydose = TRUE,
  yproj = TRUE,
  yproj_xpos = 0,
```

```

    yproj_dodge = 0.2,
    yaxis_position = c("left", "right"),
    facet_formula = NULL,
    theme_certara = TRUE,
    return_list = FALSE
)

```

Arguments

data	Data to use with multiple endpoints stacked into response (values), Endpoint(endpoint name)
response	name of the column holding the response values
endpoint	name of the column holding the name/key of the endpoint default to Endpoint
model_type	type of the trend fit one of "logistic", "none"
DOSE	name of the column holding the DOSE values default to DOSE
color_fill	name of the column to be used for color/fill default to DOSE column
logistic_by_color_fill	fit split by color? default FALSE
exposure_metrics	<p>name(s) of the column(s) to be stacked into expname exptile and split into exposure_metric_split</p>
exposure_metric_split	one of "median", "tertile", "quartile", "none"
exposure_metric_soc_value	special exposure code for standard of care default -99
exposure_metric_plac_value	special exposure code for placebo default 0
exposure_distribution	one of distributions, lineranges, boxplots or none
exposure_distribution_percent	show percent of distribution between binlimits TRUE/FALSE
exposure_distribution_percent_text_size	distribution percentages text size default to 5
dose_plac_value	string identifying placebo in DOSE column
xlab	text to be used as x axis label
ylab	text to be used as y axis label
points_alpha	alpha transparency for points
points_show	show the observations TRUE/FALSE
prob_obs_byexptile	observed probability by exptile TRUE/FALSE
prob_obs_byexptile_plac	observed probability by exptile placebo TRUE/FALSE

```

prob_obs_byexptile_group
    additional grouping for exptile probabilities default none
prob_text_size  probability text size default to 5
prob_obs_bydose
    observed probability by dose TRUE/FALSE
prob_obs_bydose_plac
    observed probability by placebo dose TRUE/FALSE
Nresp_Ntot_show
    show N responders/Ntotal ? TRUE/FALSE
Nresp_Ntot_size
    N responders/Ntotal text size default to 5
Nresp_Ntot_ypos
    y position for N responders/Ntotal two text elements the first for by exptile and
    the second for by dose/color options include with percentages top bottom
Nresp_Ntot_sep character string to separate N responders/ Ntotal default /
binlimits_show show the binlimits vertical lines TRUE/FALSE
binlimits_text_size
    binlimits text size default to 5
binlimits_ypos binlimits y position default to 0
binlimits_color
    binlimits text color default to alpha("gray70",0.5)
dist_position_scaler
    space occupied by the distribution default to 0.2
dist_offset offset where the distribution position starts default to 0
dist_scale scaling parameter for ggridges default to 0.9
lineranges_ypos
    where to put the lineranges -1
lineranges_dodge
    lineranges vertical dodge value 1
lineranges_doselabel
    TRUE/FALSE
proj_bydose project the probabilities on logistic curve TRUE/FALSE
yproj project the probabilities on y axis TRUE/FALSE
yproj_xpos y projection x position 0
yproj_dodge y projection dodge value 0.2
yaxis_position where to put y axis "left" or "right"
facet_formula facet formula to be use otherwise endpoint ~ expname
theme_certara apply certara colors and format for strips and default colour/fill
return_list What to return if True a list of the datasets and plot is returned instead of only
    the plot

```

Examples

```

# Example 1
library(ggplot2)
effICGI <- logistic_data |>
  dplyr::filter(!is.na(ICGI))|>
  dplyr::filter(!is.na(AUC))
effICGI$DOSE <- factor(effICGI$DOSE,
                        levels=c("0", "600", "1200","1800","2400"),
                        labels=c("Placebo", "600 mg", "1200 mg","1800 mg","2400 mg"))
effICGI$STUDY <- factor(effICGI$STUDY)
effICGI$ICGI2 <- effICGI$ICGI
effICGI <- tidyr::gather(effICGI,Endpoint,response,ICGI,ICGI2)
gglogisticexpdist(data = effICGI |>
  dplyr::filter(Endpoint=="ICGI"),
  response = "response",
  endpoint = "Endpoint",
  exposure_metrics = c("AUC"),
  exposure_metric_split = c("quartile"),
  exposure_metric_soc_value = -99,
  exposure_metric_plac_value = 0,
  exposure_distribution ="distributions",
  yproj_xpos = -15,
  yproj_dodge = 10,
  dist_position_scaler = 0.1,
  dist_offset = -0.1,
  Nresp_Ntot_ypos = c("with percentages","bottom"),
  prob_obs_bydose_plac = TRUE,
  prob_obs_byexptile_plac = FALSE,
  prob_obs_byexptile_group = "none",
  binlimits_ypos = 0.3,
  binlimits_color = "#475c6b",
  points_alpha= 0.8)

# Example 2
gglogisticexpdist(data = effICGI |>
  dplyr::filter(Endpoint=="ICGI"),
  response = "response",
  endpoint = "Endpoint",
  exposure_metrics = c("CMAX"),
  exposure_metric_split = c("tertile"),
  exposure_metric_soc_value = -99,
  exposure_metric_plac_value = 0,
  exposure_distribution ="lineranges",
  lineranges_ypos = -0.2,
  lineranges_dodge = 0.2,
  lineranges_doselabel = TRUE,
  Nresp_Ntot_ypos = c("with percentages","bottom"),
  prob_obs_bydose_plac = TRUE,
  prob_obs_byexptile_plac = FALSE,
  prob_obs_byexptile_group = "none", yproj_xpos = -1,
  yproj_dodge = 2,
  binlimits_color = "#475c6b",
  points_alpha= 0.8)
```

```

    dist_position_scaler = 0.1)

## Not run:
#' # Example 3
library(ggh4x)
gglogisticexpdist(data = effICGI |>
  dplyr::filter(Endpoint=="ICGI"),
  response = "response",
  endpoint = "Endpoint",
  DOSE = "DOSE",
  exposure_metrics = c("AUC"),
  exposure_metric_split = c("quartile"),
  exposure_distribution ="distributions",
  exposure_metric_soc_value = -99,
  exposure_metric_plac_value = 0,
  dist_position_scaler = 0.15)+
  facet_grid2(Endpoint~expname+DOSE2,scales="free",
  margins = "DOSE2",strip = strip_nested())
# Example 4
effICGI$SEX <- as.factor(effICGI$SEX)
gglogisticexpdist(data = effICGI |>
  dplyr::filter(Endpoint=="ICGI"),
  response = "response",
  endpoint = "Endpoint",
  DOSE = "DOSE",
  color_fill = "SEX",
  exposure_metrics = c("AUC"),
  exposure_metric_split = c("quartile"),
  exposure_distribution ="distributions",
  exposure_metric_soc_value = -99,
  exposure_metric_plac_value = 0,
  lineranges_ypos = -0.2,
  yproj_xpos = -10,
  yproj_dodge = 20,
  prob_text_size = 6,
  binlimits_text_size = 6,
  Nresp_Ntot_show = TRUE,
  dist_position_scaler = 0.15)+
  ggplot2::scale_x_continuous(breaks = seq(0,350,50),
  expand = ggplot2::expansion(add= c(0,0),mult=c(0,0)))+
  ggplot2::coord_cartesian(xlim = c(-30,355))+ 
  ggplot2::facet_grid(Endpoint~expname+color_fill2, margins ="color_fill2" )

#Example 4b
effICGI$SEX <- as.factor(effICGI$SEX)
gglogisticexpdist(data = effICGI |>
  dplyr::filter(Endpoint =="ICGI"),
  response = "response",
  endpoint = "Endpoint",
  color_fill = "SEX",
  exposure_metrics = c("AUC"),
  exposure_metric_split = c("quartile"),

```

```

exposure_metric_soc_value = -99,
exposure_metric_plac_value = 0,
dist_position_scaler = 1, dist_offset = -1 ,
yproj_xpos = -20 ,
yproj_dodge = 20 ,
exposure_distribution ="lineranges",
lineranges_doselabel = TRUE)

#Example 5
gglogisticexpdist(data = effICGI |> dplyr::filter(Endpoint=="ICGI"),
                    response = "response",
                    endpoint = "Endpoint",
                    DOSE = "DOSE",
                    exposure_metrics = c("AUC"),
                    exposure_metric_split = c("quartile"),
                    exposure_distribution ="distributions",
                    exposure_metric_soc_value = -99,
                    exposure_metric_plac_value = 0,
                    dist_position_scaler = 0.15) +
  facet_grid(Endpoint~expname+exptile,scales="free",
             margins = "exptile")

#Example 6
a <- gglogisticexpdist(data = effICGI, #
                        response = "response",
                        endpoint = "Endpoint",
                        DOSE = "DOSE",yproj_dodge = 36,
                        exposure_metrics = c("AUC"),
                        exposure_metric_split = c("quartile"),
                        exposure_distribution ="lineranges",
                        exposure_metric_soc_value = -99,
                        exposure_metric_plac_value = 0) +
  facet_grid(Endpoint~expname,switch = "both")
b <-  gglogisticexpdist(data = effICGI, #
                        response = "response",
                        endpoint = "Endpoint",
                        DOSE = "DOSE",yproj_dodge = 2,
                        exposure_metrics = c("CMAX"),
                        exposure_metric_split = c("quartile"),
                        exposure_distribution ="lineranges",
                        exposure_metric_soc_value = -99,
                        exposure_metric_plac_value = 0,
                        yaxis_position = "right") +
  facet_grid(Endpoint~expname,switch = "x")+
  theme(strip.text.y.right = element_blank(),
        strip.background.y = element_blank())
library(patchwork)
(a | b ) +
  plot_layout(guides = "collect", axes = "collect_x")&
  theme(legend.position = "top")

#Example 7
effICGI <- logistic_data |>
  dplyr::filter(!is.na(ICGI))|>

```

```

dplyr::filter(!is.na(AUC))
effICGI$DOSE <- factor(effICGI$DOSE,
                        levels=c("0", "600", "1200", "1800", "2400"),
                        labels=c("Placebo", "600 mg", "1200 mg", "1800 mg", "2400 mg"))
effICGI$STUDY <- factor(effICGI$STUDY)
effICGI$ICGI2 <- ifelse(effICGI$ICGI7 < 4,1,0)
effICGI$ICGI3 <- ifelse(effICGI$ICGI7 < 5,1,0)

effICGI <- tidyr::gather(effICGI,Endpoint,response,ICGI,ICGI2,ICGI3)
effICGI$endpointcol2 <- effICGI$Endpoint
effICGI$endpointcol3 <- effICGI$Endpoint
gglogisticexpdist(data = effICGI,
                    response = "response",
                    endpoint = "Endpoint",
                    exposure_metrics = c("AUC"),
                    exposure_metric_split = c("tertile"),
                    exposure_metric_soc_value = -99,
                    exposure_metric_plac_value = 0,
                    color_fill = "endpointcol2",
                    logistic_by_color_fill = TRUE,
                    Nresp_Ntot_show = TRUE,
                    Nresp_Ntot_ypos = c("with percentages","bottom"),
                    prob_obs_byexptile = FALSE,
                    prob_obs_byexptile_group="endpointcol3",
                    prob_obs_byexptile_plac = FALSE,
                    prob_obs_bydose_plac = TRUE,
                    binlimits_color = "#475c6b",
                    exposure_distribution ="distributions",
                    prob_obs_bydose = TRUE,
                    proj_bydose = FALSE,
                    yproj = FALSE,
                    dist_position_scaler = 0.1,
                    dist_offset = -0.1)+
  facet_grid(expname~Endpoint,scales="free_x")
#Example 8
gglogisticexpdist(data = effICGI,
                    response = "response",
                    endpoint = "endpointcol2",
                    exposure_metrics = c("AUC"),
                    exposure_metric_split = c("quartile"),
                    exposure_metric_soc_value = -99,
                    exposure_metric_plac_value = 0,
                    color_fill = "Endpoint",
                    logistic_by_color_fill = TRUE,
                    Nresp_Ntot_show = FALSE, points_show = FALSE,
                    prob_obs_byexptile = TRUE,
                    prob_obs_byexptile_group="endpointcol3",
                    prob_obs_byexptile_plac = TRUE,
                    prob_obs_bydose = FALSE,
                    prob_obs_bydose_plac = FALSE,
                    binlimits_color = "#475c6b",
                    exposure_distribution ="distributions",
                    proj_bydose = FALSE,

```

```

      yproj = FALSE,
      dist_position_scaler = 0.1,
      dist_offset = -0.1) +
  facet_grid(expname ~ ., scales = "free_x")

## End(Not run)

```

ggresponseexpdist *Create a fit vs exposure(s)plot*

Description

Produces a fit as per model_type plot with a facetable exposures/quantiles/distributions in ggplot2

Usage

```

ggresponseexpdist(
  data = dplyr::filter(logistic_data, !is.na(ICGI)),
  response = "response",
  endpoint = "Endpoint",
  model_type = c("loess", "linear", "logistic", "none"),
  DOSE = "DOSE",
  color_fill = "DOSE",
  fit_by_color_fill = FALSE,
  exposure_metrics = c("AUC", "CMAX"),
  exposure_metric_split = c("median", "tertile", "quartile", "none"),
  exposure_metric_soc_value = -99,
  exposure_metric_plac_value = 0,
  exposure_metric_soc_name = "SOC",
  exposure_metric_plac_name = "Placebo",
  exposure_distribution = c("distributions", "lineranges", "boxplots", "none"),
  exposure_distribution_percent = c("none", "%", "N (%)", "N"),
  exposure_distribution_Ntotal = c("none", "left", "right"),
  exposure_distribution_percent_text_size = 5,
  dose_plac_value = "Placebo",
  xlab = "Exposure Values",
  ylab = "Response",
  points_alpha = 0.2,
  points_show = TRUE,
  mean_obs_byexptile = TRUE,
  mean_obs_byexptile_plac = TRUE,
  mean_obs_byexptile_text_size = 5,
  mean_obs_byexptile_group = "none",
  mean_obs_bydose = FALSE,
  mean_obs_bydose_plac = FALSE,
  mean_obs_bydose_text_size = 5,
  N_byexptile_ypos = c("with means", "top", "bottom", "none"),

```

```

N_bydose_ypos = c("with means", "top", "bottom", "none"),
N_text_size = 5,
N_text_sep = NULL,
binlimits_show = TRUE,
binlimits_text_size = 5,
binlimits_ypos = 0,
binlimits_color = "#B3B3B380",
dist_position_scaler = 0.2,
dist_offset = 0,
dist_scale = 0.9,
lineranges_ypos = NULL,
lineranges_dodge = NULL,
lineranges_doselabel = FALSE,
lineranges_Ntotal = c("none", "left", "right"),
proj_bydose = FALSE,
yproj = FALSE,
yproj_xpos = 0,
yproj_dodge = 0.2,
yaxis_position = c("left", "right"),
facet_formula = NULL,
theme_certara = TRUE,
color_legend_title = "",
fill_legend_title = "",
linetype_legend_title = "",
shape_legend_title = "",
combine_fill_linetype_legend = TRUE,
legend_order = c("model", "color", "linetype", "shape"),
return_list = FALSE
)

```

Arguments

<code>data</code>	Data to use with multiple endpoints stacked into response (values), Endpoint(endpoint name)
<code>response</code>	name of the column holding the response values
<code>endpoint</code>	name of the column holding the name/key of the endpoint default to Endpoint
<code>model_type</code>	type of the trend fit one of "loess", "linear", "logistic", "none"
<code>DOSE</code>	name of the column holding the DOSE/regimen values default to DOSE should be a factor
<code>color_fill</code>	name of the column to be used for color/fill default to DOSE column should be a factor
<code>fit_by_color_fill</code>	split fit by color/fill? default FALSE
<code>exposure_metrics</code>	name(s) of the column(s) to be stacked into expname exptile and split into exposure_metric_split

```

exposure_metric_split
    one of "median", "tertile", "quartile", "none"
exposure_metric_soc_value
    special exposure code for standard of care default -99
exposure_metric_plac_value
    special exposure code for placebo default 0
exposure_metric_soc_name
    soc name default to "soc"
exposure_metric_plac_name
    placebo name default to "placebo"
exposure_distribution
    one of distributions, lineranges, boxplots or none
exposure_distribution_percent
    show N/percent of distribution between binlimits one of "%", "N (%)", "N", "none"
exposure_distribution_Ntotal
    show Ntotal by dose level next to the distribution one of "left", "right", "none"
exposure_distribution_percent_text_size
    distribution percentages text size default to 5
dose_plac_value
    string identifying placebo in DOSE column
xlab
    text to be used as x axis label
ylab
    text to be used as y axis label
points_alpha
    alpha transparency for points
points_show
    show the observations TRUE/FALSE
mean_obs_byexptile
    observed mean by exptile TRUE/FALSE
mean_obs_byexptile_plac
    observed mean by exptile placebo TRUE/FALSE
mean_obs_byexptile_text_size
    by exptile mean text size default to 5
mean_obs_byexptile_group
    additional grouping for exptile means default none
mean_obs_bydose
    observed mean by dose TRUE/FALSE
mean_obs_bydose_plac
    observed mean by placebo dose TRUE/FALSE
mean_obs_bydose_text_size
    by dose mean text size default to 5
N_byexptile_ypos
    N responders/Ntotal y position by exptile one of with means top bottom none
N_bydose_ypos
    N responders/Ntotal y position by dose/color one of with means top bottom
    none
N_text_size
    N responders/Ntotal text size default to 5

```

N_text_sep character string to separate N responders/Ntotal or N/mean defaults to / otherwise \n

binlimits_show show the binlimits vertical lines TRUE/FALSE

binlimits_text_size
binlimits text size default to 5

binlimits_ypos binlimits y position default to -Inf

binlimits_color
binlimits text color default to alpha("gray70",0.5)

dist_position_scaler
space occupied by the distribution default to 0.2

dist_offset offset where the distribution position starts default to 0

dist_scale scaling parameter for ggridges default to 0.9

lineranges_ypos
where to put the lineranges -1

lineranges_dodge
lineranges vertical dodge value 1

lineranges_doselabel
TRUE/FALSE

lineranges_Ntotal
show Ntotal by dose level next to the lineranges one of "left","right","none"

proj_bydose project the predictions on the fit curve TRUE/FALSE

yproj project the predictions on y axis TRUE/FALSE

yproj_xpos y projection x position 0

yproj_dodge y projection dodge value 0.2

yaxis_position where to put y axis "left" or "right"

facet_formula facet formula to be use otherwise endpoint ~ expname

theme_certara apply certara colors and format for strips and default colour/fill

color_legend_title
text for colour legend title

fill_legend_title
text for fill legend title

linetype_legend_title
text for linetype legend title

shape_legend_title
text for shape legend title

combine_fill_linetype_legend
defaults to true

legend_order Legend order. A four-element vector with the following items ordered in your desired order: "model", "color", "linetype", "shape". if an item is absent the legend will be omitted.

return_list What to return if True a list of the datasets and plot is returned instead of only the plot

Examples

```

# Example 1
library(ggplot2)
effICGI <- logistic_data |>
dplyr::filter(!is.na(ICGI))|>
dplyr::filter(!is.na(AUC))
effICGI$DOSE <- factor(effICGI$DOSE,
levels=c("0", "600", "1200","1800","2400"),
labels=c("Placebo", "600 mg", "1200 mg","1800 mg","2400 mg"))
effICGI$STUDY <- factor(effICGI$STUDY)
effICGI$ICGI2 <- effICGI$ICGI
effICGI <- tidyr::gather(effICGI,Endpoint,response,ICGI,ICGI2)

ggresponseexpdist(data = effICGI |>
dplyr::filter(Endpoint=="ICGI"),
model_type = "loess",
exposure_metrics = c("AUC","CMAX"),
legend_order = c("color","shape","model"),
color_legend_title ="Dose\nLevels")

# Example 2
ggresponseexpdist(data = effICGI |>
dplyr::filter(Endpoint=="ICGI"),
model_type = "logistic",
exposure_metrics = c("AUC","CMAX"),
exposure_distribution ="boxplots")

# Example 3
ggresponseexpdist(data = effICGI|>
dplyr::filter(Endpoint=="ICGI"),
model_type = "linear",
exposure_metrics = c("AUC","WT"),
exposure_distribution ="lineranges")

# Example 4
ggresponseexpdist(data = effICGI |>
dplyr::filter(Endpoint=="ICGI"),
response = "response",
endpoint = "Endpoint",
model_type = "loess",
DOSE = "DOSE",
color_fill = "DOSE",
exposure_metrics = c("AUC","CMAX"),
exposure_metric_split = c("tertile"),
exposure_metric_soc_value = -99,
exposure_metric_plac_value = 0,
exposure_distribution ="distributions",
exposure_distribution_Ntotal="right",
N_byexptile_ypos = "top",
mean_obs_bydose = TRUE,
mean_obs_bydose_text_size = 0,
mean_obs_byexptile_text_size = 5,
```

```

N_bydose_ypos = "none",
N_text_sep = "/",
binlimits_color = "#475c6b",
binlimits_ypos = 0.2,
points_alpha= 0.1)

## Not run:
# Example 5
effICGI <- logistic_data |>
dplyr::filter(!is.na(ICGI))|>
dplyr::filter(!is.na(AUC))
effICGI$DOSE <- factor(effICGI$DOSE,
levels=c("0", "600", "1200","1800","2400"),
labels=c("Placebo", "600 mg", "1200 mg","1800 mg","2400 mg"))
effICGI$STUDY <- factor(effICGI$STUDY)
effICGI$ICGI2 <- ifelse(effICGI$ICGI7 < 4,1,0)
effICGI$ICGI3 <- ifelse(effICGI$ICGI7 < 5,1,0)

effICGI <- tidyr::gather(effICGI,Endpoint,response,ICGI,ICGI2,ICGI3)
effICGI$endpointcol2 <- effICGI$Endpoint
effICGI$endpointcol3 <- effICGI$Endpoint

ggresponseexpdist(data = effICGI,
points_show = FALSE,
exposure_metrics = c("AUC"),
exposure_distribution ="lineranges",
color_fill = "endpointcol2",
model_type = "logistic",
fit_by_color_fill = TRUE,
mean_obs_byexptile_text_size = 0,
mean_obs_byexptile_group="endpointcol3",
facet_formula = Endpoint~expname,
N_byexptile_ypos = "none",
binlimits_text_size = 0
)+

ggplot2::facet_grid(expname~Endpoint,margin="Endpoint")

# Example 6 retrun a list and customize
plist <- ggresponseexpdist(data = effICGI |>
dplyr::filter(Endpoint=="ICGI"),
model_type = "logistic",
mean_obs_byexptile = TRUE,
N_byexptile_ypos = "none",
mean_obs_byexptile_text_size = 4,
binlimits_ypos = -Inf,
exposure_metric_split = "tertile",
exposure_metrics = c("AUC"),
return_list = TRUE)
byexptileinformation <- plist[[7]]
plotwihoutlabels <- plist[[9]]

#construct text label
byexptileinformation <- byexptileinformation %>%

```

```

dplyr::group_by(exptile,expname)%>%
dplyr::mutate(meanbin = mean(c(minexp,maxexp))) %>%
dplyr::mutate(label = ifelse(exptile!="Placebo",
                           paste0(exptile,"\\n","[",round(minexp,0),"-",round(maxexp,0),"]",
                           "\\n",N,"\\n",Ntot),
                           paste0(exptile,"\\n","","\\n",N,"\\n",Ntot)),
                           x_pos = ifelse(exptile=="Placebo",-25,meanbin )
)
plotwithoutlabels +
  geom_text(data=byexptileinformation,size = 3,
            aes(x=x_pos ,label=label,y = 0.5),
            inherit.aes = FALSE,
            vjust= 1, hjust = 0.5) +
  geom_text(data=data.frame(
    label="exposure ntile\\n[min-max]\\nN responders\\nN Total"),size = 3,
            x=Inf,y = 0.5,
            aes(label=label),inherit.aes = FALSE,
            vjust= 1, hjust = 1) +
  facet_wrap(~expname,ncol=2,scales = "free_x")

## End(Not run)

```

logistic_data*Simulated Exposure Response Data***Description**

A dataset containing data suitable for logistic regression

Usage

```
logistic_data
```

Format

A data frame with 600 rows and 10 variables

STUDY Study identifier

ID Subject Identifier

DOSE Dose, in mg

GBDS Dose, in alternative salt

SEX Sex of the subject

AGE age of the subject, in years

WT weight of the subject, in kg

RACE Race of the subject

CRCL Creatinine clearance
BRLS RLS score
PRLS RLS score
AUC Area under the curve exposure
CMAX Maximun concentration exposure
ICGI response 0/1
ICGI7 response 1 to 7

Source

inspired from a real data submission

Examples

```
logistic_data
```

<code>run_ggquickeda</code>	<i>Run the ggquickeda application</i>
-----------------------------	---------------------------------------

Description

Run the ggquickeda application.

Usage

```
run_ggquickeda(data = NULL, ...)
```

Arguments

<code>data</code>	The initial data.frame to load into the application.
<code>...</code>	Additional arguments for bookmarking

Examples

```
if (interactive()) {
  run_ggquickeda()
}
```

sample_data

Simulated Pharmacokinetic Concentration Data

Description

A dataset containing concentration-time data with the given dose and some subject characteristics to help in the app exploration.

Usage

```
sample_data
```

Format

A data frame with 600 rows and 10 variables

ID Subject Identifier, an integer from 1 to 150

Time Time of dose given or drug sample measured, in hours

Amt dose given at the corresponding Time, in milligrams

Conc drug concentrations in the plasma sample, in mg/L

Age age of the subject, in years

Weight weight of the subject, in kg

Gender Sex of the subject, a factor with Female and Male levels

Race Race of the subject, a factor with Asian, Black, Caucasian, Hispanic and Other levels

Dose dose group of the subject, in milligrams

AGECAT age category of the subject, a variable cutting Age into two values 0/1

Source

"sd_oral_richpk" from 'PKPDmisc' R package with an additional AGECAT variable

Examples

```
sample_data
```

stat_km*Adds a Kaplan Meier Estimate of Survival*

Description

Adds a Kaplan Meier Estimate of Survival

Usage

```
stat_km(
  mapping = NULL,
  data = NULL,
  geom = "km",
  position = "identity",
  show.legend = NA,
  inherit.aes = TRUE,
  trans = scales::identity_trans(),
  firstx = 0,
  firsty = 1,
  type = "kaplan-meier",
  start.time = 0,
  ...
)
```

Arguments

mapping	Set of aesthetic mappings created by aes() . If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply <code>mapping</code> if there is no plot mapping.
data	The data to be displayed in this layer. There are three options: If <code>NULL</code> , the default, the data is inherited from the plot data as specified in the call to ggplot() . A <code>data.frame</code> , 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 <code>data.frame</code> , and will be used as the layer data. A function can be created from a formula (e.g. <code>~ head(.x, 10)</code>).
geom	The geometric object to use to display the data for this layer. When using a <code>stat_*</code> function to construct a layer, the <code>geom</code> argument can be used to override the default coupling between stats and geoms. The <code>geom</code> argument accepts the following: <ul style="list-style-type: none"> • A Geom ggproto subclass, for example <code>GeomPoint</code>. • A string naming the geom. To give the geom as a string, strip the function name of the <code>geom_</code> prefix. For example, to use <code>geom_point()</code>, give the geom as "point".

	<ul style="list-style-type: none"> For more information and other ways to specify the geom, see the layer geom 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: <ul style="list-style-type: none"> The result of calling a position function, such as <code>position_jitter()</code>. 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 <code>position_</code> prefix. For example, to use <code>position_jitter()</code>, give the position as "jitter". For more information and other ways to specify the position, see the layer position documentation.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display. To include legend keys for all levels, even when no data exists, use TRUE. If NA, all levels are shown in legend, but unobserved levels are omitted.
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. annotation_borders() .
trans	Transformation to apply to the survival probabilities. Defaults to "identity". Other options include "event", "cumhaz", "cloglog", or define your own using trans_new .
firstx, firsty	the starting point for the survival curves. By default, the plot program obeys tradition by having the plot start at (0,1).
type	an older argument that combined stype and ctype, now deprecated. Legal values were "kaplan-meier" which is equivalent to <code>stype=1, ctype=1</code> , "fleming-harrington" which is equivalent to <code>stype=2, ctype=1</code> , and "fh2" which is equivalent to <code>stype=2, ctype=2</code> .
start.time	numeric value specifying a time to start calculating survival information. The resulting curve is the survival conditional on surviving to start.time.
...	Other arguments passed to survfit.formula

Details

This stat is for computing the confidence intervals for the Kaplan-Meier survival estimate for right-censored data. It requires the aesthetic mapping `x` for the observation times and `status` which indicates the event status, 0=alive, 1=dead or 1/2 (2=death). Logical status is not supported.

Value

a data.frame with additional columns:

x	x in data
y	Kaplan-Meier Survival Estimate at x

Aesthetics

`stat_km` understands the following aesthetics (required aesthetics are in bold):

- **time** The survival times
- **status** The censoring indicator, see [Surv](#) for more information.
- **alpha**
- **color**
- **linetype**
- **size**

Examples

```
library(ggplot2)
sex <- rbinom(250, 1, .5)
df <- data.frame(time = exp(rnorm(250, mean = sex)), status = rbinom(250, 1, .75), sex = sex)
ggplot(df, aes(time = time, status = status, color = factor(sex))) +
  stat_km()

## Examples illustrating the options passed to survfit.formula

p1 <- ggplot(df, aes(time = time, status = status))
p1 + stat_km()
p1 + stat_km(trans = "cumhaz")
# for cloglog plots also log transform the time axis
p1 + stat_km(trans = "cloglog") + scale_x_log10()
p1 + stat_km(type = "fleming-harrington")
p1 + stat_km(start.time = 5)
```

`stat_kmband`

Adds confidence bands to a Kaplan Meier Estimate of Survival

Description

Adds confidence bands to a Kaplan Meier Estimate of Survival

Usage

```
stat_kmband(
  mapping = NULL,
  data = NULL,
  geom = "kmband",
  position = "identity",
  show.legend = NA,
  inherit.aes = TRUE,
  trans = "identity",
  firstx = 0,
```

```

firsty = 1,
type = "kaplan-meier",
error = "greenwood",
conf.type = "log",
conf.lower = "usual",
start.time = 0,
conf.int = 0.95,
...
)

```

Arguments

<code>mapping</code>	Set of aesthetic mappings created by aes() . If specified and <code>inherit.aes</code> = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply <code>mapping</code> if there is no plot mapping.
<code>data</code>	The data to be displayed in this layer. There are three options: If <code>NULL</code> , the default, the data is inherited from the plot data as specified in the call to ggplot() . A <code>data.frame</code> , 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 <code>data.frame</code> , and will be used as the layer data. A function can be created from a <code>formula</code> (e.g. <code>~ head(.x, 10)</code>).
<code>geom</code>	The geometric object to use to display the data for this layer. When using a <code>stat_*</code> function to construct a layer, the <code>geom</code> argument can be used to override the default coupling between stats and geoms. The <code>geom</code> argument accepts the following: <ul style="list-style-type: none"> • A Geom ggproto subclass, for example <code>GeomPoint</code>. • A string naming the geom. To give the geom as a string, strip the function name of the <code>geom_</code> prefix. For example, to use <code>geom_point()</code>, give the geom as "point". • For more information and other ways to specify the geom, see the layer geom documentation.
<code>position</code>	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 <code>position</code> argument accepts the following: <ul style="list-style-type: none"> • The result of calling a position function, such as <code>position_jitter()</code>. 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 <code>position_</code> prefix. For example, to use <code>position_jitter()</code>, give the position as "jitter". • For more information and other ways to specify the position, see the layer position documentation.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It

	can also be a named logical vector to finely select the aesthetics to display. To include legend keys for all levels, even when no data exists, use TRUE. If NA, all levels are shown in legend, but unobserved levels are omitted.
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. annotation_borders() .
trans	Transformation to apply to the survival probabilities. Defaults to "identity". Other options include "event", "cumhaz", "cloglog", or define your own using scales::trans_new() .
firstx, firsty	the starting point for the survival curves. By default, the plot program obeys tradition by having the plot start at (0,1).
type	an older argument that combined stype and ctype, now deprecated. Legal values were "kaplan-meier" which is equivalent to stype=1, ctype=1, "fleming-harrington" which is equivalent to stype=2, ctype=1, and "fh2" which is equivalent to stype=2, ctype=2.
error	either the string "greenwood" for the Greenwood formula or "tsiatis" for the Tsiatis formula, (only the first character is necessary). The default is "greenwood".
conf.type	One of "none", "plain", "log" (the default), "log-log" or "logit".
conf.lower	a character string to specify modified lower limits to the curve, the upper limit remains unchanged. Possible values are "usual" (unmodified), "peto", and "modified". The modified lower limit is based on an "effective n" argument. The confidence bands will agree with the usual calculation at each death time, but unlike the usual bands the confidence interval becomes wider at each censored observation. The extra width is obtained by multiplying the usual variance by a factor m/n, where n is the number currently at risk and m is the number at risk at the last death time. (The bands thus agree with the un-modified bands at each death time.) This is especially useful for survival curves with a long flat tail. The Peto lower limit is based on the same "effective n" argument as the modified limit, but also replaces the usual Greenwood variance term with a simple approximation. It is known to be conservative.
start.time	numeric value specifying a time to start calculating survival information. The resulting curve is the survival conditional on surviving to start.time.
conf.int	the level for a two-sided confidence interval on the survival curve(s). Default is 0.95.
...	Other arguments passed to survfit.formula

Details

This stat is for computing the confidence intervals for the Kaplan-Meier survival estimate for right-censored data. It requires the aesthetic mapping `x` for the observation times and `status` which indicates the event status, 0=alive, 1=dead or 1/2 (2=death). Logical status is not supported.

Value

a data.frame with additional columns:

x	x in data
ymin	Lower confidence limit of KM curve
ymax	Upper confidence limit of KM curve

Aesthetics

stat_kmband understands the following aesthetics (required aesthetics are in bold):

- time The survival times
- status The censoring indicator, see [Surv](#) for more information.
- alpha
- color
- linetype
- linewidth

Examples

```
library(ggplot2)
sex <- rbinom(250, 1, .5)
df <- data.frame(time = exp(rnorm(250, mean = sex)), status = rbinom(250, 1, .75), sex = sex)
ggplot(df, aes(time = time, status = status, color = factor(sex))) +
  stat_km()

## Examples illustrating the options passed to survfit.formula

p1 <- ggplot(df, aes(time = time, status = status))
p1 + stat_km() + stat_kmband(conf.int = .99)
p1 + stat_kmband(error = "greenwood", fill="red",alpha=0.2) +
  stat_kmband(error = "tsiatis", fill="blue",alpha=0.2)+ stat_km()
p1 + stat_km() + stat_kmband(conf.type = "log-log")+ stat_kmband(conf.type = "log")
```

stat_kmticks

Adds tick marks to a Kaplan Meier Estimate of Survival

Description

Adds tick marks to a Kaplan Meier Estimate of Survival

Usage

```
stat_kmticks(
  mapping = NULL,
  data = NULL,
  geom = "kmticks",
  position = "identity",
  show.legend = NA,
```

```
inherit.aes = TRUE,
trans,
...
)
```

Arguments

<code>mapping</code>	Set of aesthetic mappings created by aes() . If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply <code>mapping</code> if there is no plot mapping.
<code>data</code>	The data to be displayed in this layer. There are three options: If <code>NULL</code> , the default, the data is inherited from the plot data as specified in the call to ggplot() . A <code>data.frame</code> , 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 <code>data.frame</code> , and will be used as the layer data. A function can be created from a formula (e.g. <code>~ head(.x, 10)</code>).
<code>geom</code>	The geometric object to use to display the data for this layer. When using a <code>stat_*</code> function to construct a layer, the <code>geom</code> argument can be used to override the default coupling between stats and geoms. The <code>geom</code> argument accepts the following: <ul style="list-style-type: none"> • A Geom ggproto subclass, for example <code>GeomPoint</code>. • A string naming the geom. To give the geom as a string, strip the function name of the <code>geom_</code> prefix. For example, to use <code>geom_point()</code>, give the geom as "point". • For more information and other ways to specify the geom, see the layer geom documentation.
<code>position</code>	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 <code>position</code> argument accepts the following: <ul style="list-style-type: none"> • The result of calling a position function, such as <code>position_jitter()</code>. 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 <code>position_</code> prefix. For example, to use <code>position_jitter()</code>, give the position as "jitter". • For more information and other ways to specify the position, see the layer position documentation.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes. It can also be a named logical vector to finely select the aesthetics to display. To include legend keys for all levels, even when no data exists, use TRUE. If NA, all levels are shown in legend, but unobserved levels are omitted.
<code>inherit.aes</code>	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. annotation_borders() .

trans	Transformation to apply to the survival probabilities. Defaults to "identity". Other options include "event", "cumhaz", "cloglog", or define your own using trans_new
...	Other arguments passed to survfit.formula

Details

This stat is for computing the tick marks for a Kaplan-Meier survival estimate for right-censored data. The tick marks will appear at each censoring time which is also not a death time, which is the default for [plot.survfit](#). It requires the aesthetic mapping `x` for the observation times and `status` which indicates the event status, normally 0=alive, 1=dead. Other choices are TRUE/FALSE (TRUE = death) or 1/2 (2=death).

Value

a data.frame with additional columns:

x	x in data
y	Kaplan-Meier Survival Estimate at x

Aesthetics

`stat_kmticks` understands the following aesthetics (required aesthetics are in bold):

- `time` The survival times
- `status` The censoring indicator, see [Surv](#) for more information.
- `alpha`
- `color`
- `linetype`
- `size`

See Also

[stat_km](#); [stat_kmband](#)

Examples

```
library(ggplot2)
sex <- rbinom(250, 1, .5)
df <- data.frame(time = exp(rnorm(250, mean = sex)), status = rbinom(250, 1, .75), sex = sex)
ggplot(df, aes(time = time, status = status, color = factor(sex))) +
  stat_km() + stat_kmticks()
```

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