

Package ‘nullabor’

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Description Tools for visual inference. Generate null data sets and null plots using permutation and simulation. Calculate distance metrics for a lineup, and examine the distributions of metrics.

Title Tools for Graphical Inference

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URL <https://github.com/dicook/nullabor>

BugReports <https://github.com/dicook/nullabor/issues>

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 aud

Conversion rate of 1 Australian Doller (AUD) to 1 US Dollar

Description

The dataset consists of the daily exchange rates of 1 Australian Dollar to 1 US Dollar between Jan 9 2018 and Feb 21 2018.

bin_dist	<i>Binned Distance</i>
----------	------------------------

Description

Data X is binned into X.bin bins in x-direction and Y.bin bins in y-direction. The number of points in each cell is then counted. Same is done for data PX. An euclidean distance is calculated between the number of points in each cell between X and PX.

Usage

```
bin_dist(X, PX, lineup.dat = lineup.dat, X.bin = 5, Y.bin = 5)
```

Arguments

X	a data.frame with two variables, the first two columns are used
PX	another data.frame with two variables, the first two columns are used
lineup.dat	lineup data so that the binning is done based on the lineup data and not the individual plots, by default lineup.dat = lineup.dat ; if one wishes to calculate the binned distance between two plots, one should use lineup.dat = NULL
X.bin	number of bins on the x-direction, by default X.bin = 5
Y.bin	number of bins on the y-direction, by default Y.bin = 5

Value

distance between X and PX

Examples

```
with(mtcars, bin_dist(data.frame(wt, mpg), data.frame(sample(wt), mpg),
  lineup.dat = NULL))
```

box_dist	<i>Distance based on side by side Boxplots</i>
----------	--

Description

Assuming that data set X consists of a categorical group variable a numeric value, a summary of the first quartile, median and third quartile of this value is calculated for each group. The extent (as absolute difference) of the minimum and maximum value across groups is computed for first quartile, median and third quartile. Same is done for data PX. Finally an euclidean distance is calculated between the absolute differences of X and PX.

Usage

```
box_dist(X, PX)
```

Arguments

X a data.frame with one factor variable and one continuous variable
 PX a data.frame with one factor variable and one continuous variable

Value

distance between X and PX

Examples

```
if(require('dplyr')) {
  with(mtcars,
    box_dist(data.frame(as.factor(am), mpg),
              data.frame(as.factor(sample(am)), mpg))
  )
}
```

calc_diff	<i>Calculating the difference between true plot and the null plot with the maximum distance.</i>
-----------	--

Description

Distance metric is used to calculate the mean distance between the true plot and all the null plots in a lineup. The difference between the mean distance of the true plot and the maximum mean distance of the null plots is calculated.

Usage

```
calc_diff(lineup.dat, var, met, pos, dist.arg = NULL, m = 20)
```

Arguments

lineup.dat lineup data to get the lineup
 var a vector of names of the variables to be used to calculate the difference
 met distance metric needed to calculate the distance as a character
 pos position of the true plot in the lineup
 dist.arg a list or vector of inputs for the distance metric met; NULL by default
 m number of plots in the lineup, by default m = 20

Value

difference between the mean distance of the true plot and the maximum mean distance of the null plots

Examples

```

if(require('dplyr')){
  lineup.dat <- lineup(null_permute('mpg'), mtcars, pos = 1)
  calc_diff(lineup.dat, var = c('mpg', 'wt'), met = 'bin_dist',
  dist.arg = list(lineup.dat = lineup.dat, X.bin = 5, Y.bin = 5), pos = 1, m = 8)}

if(require('dplyr')){
  calc_diff(lineup(null_permute('mpg'), mtcars, pos = 1), var = c('mpg', 'wt'), met = 'reg_dist',
  dist.arg = NULL, pos = 1, m = 8)}

```

calc_mean_dist	<i>Calculating the mean distances of each plot in the lineup.</i>
----------------	---

Description

Distance metric is used to calculate the mean distance between the true plot and all the null plots in a lineup. The mean distances of each null plot to all the other null plots are calculated. The mean distances are returned for all the plots in the lineup.

Usage

```
calc_mean_dist(lineup.dat, var, met, pos, dist.arg = NULL, m = 20)
```

Arguments

lineup.dat	lineup data of the lineup
var	a vector of names of the variables to be used to calculate the mean distances
met	distance metric needed to calculate the distance as a character
pos	position of the true plot in the lineup
dist.arg	a list or vector of inputs for the distance metric met; NULL by default
m	number of plots in the lineup, by default m = 20

Value

the mean distances of each plot in the lineup

Examples

```

if(require('dplyr')){
  calc_mean_dist(lineup(null_permute('mpg'), mtcars, pos = 1), var = c('mpg', 'wt'),
  met = 'reg_dist', pos = 1, m = 10)}

```

 decrypt

Use decrypt to reveal the position of the real data.

Description

The real data position is encrypted by the lineup function, and writes this out as a text string. Decrypt, decrypts this text string to reveal which where the real data is.

Usage

```
decrypt(...)
```

Arguments

```
...          character vector to decrypt
```

Examples

```
decrypt('0uXR2p rut L202')
```

 distmet

Empirical distribution of the distance

Description

The empirical distribution of the distance measures is calculated based on the mean distance of each of the null plots from the other null plots in a lineup. At this moment this method works only for [null_permute](#) method. This function helps get some assessment of whether the actual data plot is very different from the null plots.

Usage

```
distmet(
  lineup.dat,
  var,
  met,
  method,
  pos,
  repl = 1000,
  dist.arg = NULL,
  m = 20
)
```

Arguments

lineup.dat	lineup data
var	a vector of names of the variables to be used
met	distance metric needed to calculate the distance as a character
method	method for generating null data sets
pos	position of the observed data in the lineup
repl	number of sets of null plots selected to obtain the distribution; 1000 by default
dist.arg	a list or vector of inputs for the distance metric met; NULL by default
m	the number of plots in the lineup; m = 20 by default

Value

lineup has the data used for the calculations

null_values contains new null samples from which to compare nulls in lineup

diff difference in distance between nulls and actual data and that of the null that is most different from other nulls. A negative value means that the actual data plot is similar to the null plots.

closest list of the five closest nulls to the actual data plot

pos position of the actual data plot in the lineup

Examples

```
# Each of these examples uses a small number of nulls (m=8), and a small number of
# repeated sampling from the null distribution (repl=100), to make it faster to run.
# In your own examples you should think about increasing each of these, at least to the defaults.
## Not run:
if (require('dplyr')) {
  d <- lineup(null_permute('mpg'), mtcars, pos = 1)
  dd <- distmet(d, var = c('mpg', 'wt'),
               'reg_dist', null_permute('mpg'), pos = 1, repl = 100, m = 8)
  distplot(dd, m=8)
}

## End(Not run)

## Not run:
d <- lineup(null_permute('mpg'), mtcars, pos=4, n=8)
library(ggplot2)
ggplot(d, aes(mpg, wt)) + geom_point() + facet_wrap(~ .sample, ncol=4)
if (require('dplyr')) {
  dd <- distmet(d, var = c('mpg', 'wt'), 'bin_dist', null_permute('mpg'),
               pos = 4, repl = 100, dist.arg = list(lineup.dat = d, X.bin = 5,
               Y.bin = 5), m = 8)
  distplot(dd, m=8)
}

## End(Not run)
```

```

# Example using bin_dist
## Not run:
if (require('dplyr')) {
  d <- lineup(null_permute('mpg'), mtcars, pos = 1)
  library(ggplot2)
  ggplot(d, aes(mpg, wt)) + geom_point() + facet_wrap(~ .sample, ncol=5)
  dd <- distmet(d, var = c('mpg', 'wt'),
    'bin_dist', null_permute('mpg'), pos = 1, repl = 500,
    dist.arg = list(lineup.dat = d, X.bin = 5, Y.bin = 5))
  distplot(dd)
}

## End(Not run)

# Example using uni_dist
## Not run:
mod <- lm(wt ~ mpg, data = mtcars)
resid.dat <- data.frame(residual = mod$resid)
d <- lineup(null_dist('residual', dist = 'normal'), resid.dat, pos=19)
ggplot(d, aes(residual)) + geom_histogram(binwidth = 0.25) + facet_wrap(~ .sample, ncol=5)
if (require('dplyr')) {
  dd <- distmet(d, var = 'residual', 'uni_dist', null_dist('residual',
    dist = 'normal'), pos = 19, repl = 500)
  distplot(dd)
}

## End(Not run)

```

distplot

Plotting the distribution of the distance measure

Description

The permutation distribution of the distance measure is plotted with the distances for the null plots. Distance measure values for the null plots and the true plot are overlaid.

Usage

```
distplot(dat, m = 20)
```

Arguments

dat	output from distmet
m	the number of plots in the lineup; m = 20 by default

Examples

```
## Not run:
if (require('dplyr')) {
  d <- lineup(null_permute('mpg'), mtcars, pos = 1)
  library(ggplot2)
  ggplot(d, aes(mpg, wt)) + geom_point() + facet_wrap(~.sample)
  distplot(distmet(d, var = c('mpg', 'wt'), 'reg_dist', null_permute('mpg'),
    pos = 1, repl = 100, m = 8), m = 8)
}

## End(Not run)
```

 electoral

Polls and election results from the 2012 US Election

Description

Polls and election results from the 2012 US Election

Format

A list with two data frames: polls is a data frame of 51 rows and 4 variables

State State name

Electoral.vote Number of electoral votes in the 2012 election

Margin Margin between the parties with the highest number of votes and second highest number of votes. These margins are based on polls.

Democrat logical vector True, if the democratic party is the majority party in this state.

election is a data frame of 51 rows and 5 variables

State State name

Candidate character string of the winner: Romney or Obama

Electoral.vote Number of electoral votes in the 2012 election

Margin Margin between the parties with the highest number of votes and second highest number of votes. These margins are based on the actual election outcome

Democrat logical vector True, if the democratic party is the majority party in this state.

 la1

Los Angeles Lakers play-by-play data.

Description

Play by play data from all games played by the Los Angeles lakers in the 2008/2009 season.

`lineup`*The line-up protocol.*

Description

In this protocol the plot of the real data is embedded amongst a field of plots of data generated to be consistent with some null hypothesis. If the observe can pick the real data as different from the others, this lends weight to the statistical significance of the structure in the plot. The protocol is described in Buja, Cook, Hofmann, Lawrence, Lee, Swayne, Wickham (2009) Statistical inference for exploratory data analysis and model diagnostics, *Phil. Trans. R. Soc. A*, 367, 4361-4383.

Usage

```
lineup(method, true = NULL, n = 20, pos = sample(n, 1), samples = NULL)
```

Arguments

<code>method</code>	method for generating null data sets
<code>true</code>	true data set. If NULL, <code>find_plot_data</code> will attempt to extract it from the current <code>ggplot2</code> plot.
<code>n</code>	total number of samples to generate (including true data)
<code>pos</code>	position of true data. Leave missing to pick position at random. Encrypted position will be printed on the command line, <code>decrypt</code> to understand.
<code>samples</code>	samples generated under the null hypothesis. Only specify this if you don't want <code>lineup</code> to generate the data for you.

Details

Generate $n - 1$ null datasets and randomly position the true data. If you pick the real data as being noticeably different, then you have formally established that it is different to with p-value $1/n$.

Examples

```
library(ggplot2)
ggplot(lineup(null_permute('mpg'), mtcars), aes(mpg, wt)) +
  geom_point() +
  facet_wrap(~ .sample)
ggplot(lineup(null_permute('cyl'), mtcars),
  aes(mpg, .sample, colour = factor(cyl))) +
  geom_point()
```

lineup_histograms	<i>Check distributional assumptions using histograms and the lineup protocol.</i>
-------------------	---

Description

This function is used to quickly create lineup plots to check distributional assumptions using histograms with kernel density estimates. The null hypothesis is that the data follows the distribution specified by the `dist` argument. In the lineup protocol the plot of the real data is embedded amongst a field of plots of data generated to be consistent with some null hypothesis. If the observer can pick the real data as different from the others, this lends weight to the statistical significance of the structure in the plot. The protocol is described in Buja et al. (2009).

Usage

```
lineup_histograms(
  data,
  variable,
  dist = NULL,
  params = NULL,
  color_bars = "black",
  fill_bars = "grey",
  color_lines = "brown3"
)
```

Arguments

<code>data</code>	a data frame.
<code>variable</code>	the name of the variable that should be plotted.
<code>dist</code>	the null distribution name. One of: "beta", "cauchy", "chi-squared", "exponential", "f", "gamma", "geometric", "log-normal", "lognormal", "logistic", "negative binomial", "binomial", "normal", "poisson", "t", "uniform", "weibull"
<code>params</code>	list of parameters of distribution. If NULL, will use <code>fitdistr</code> to estimate them if possible. For uniform, beta, and binomial distributions, the parameters must be specified. See <code>?dunif</code> , <code>?dbeta</code> , and <code>?dbinom</code> for parameter names.
<code>color_bars</code>	the color used for the borders of the bars. Can be a name or a color HEX code.
<code>fill_bars</code>	the color used to fill the bars.
<code>color_lines</code>	the color used for the density curves.

Details

19 null datasets are plotted together the the true data (randomly positioned) If you pick the real data as being noticeably different, then you have formally established that it is different to with p-value 0.05.

Run the decrypt message printed in the R Console to see which plot represents the true data.

Value

a ggplot

References

Buja, Cook, Hofmann, Lawrence, Lee, Swayne, Wickham. (2009). Statistical inference for exploratory data analysis and model diagnostics, *Phil. Trans. R. Soc. A*, 367, 4361-4383.

See Also

null_dist

Examples

```
data(tips)
lineup_histograms(tips, "total_bill", dist = "normal") # Normal distribution

# Some distributions require that the parameters be specified:
lineup_histograms(tips, "size", dist = "binomial", params = list(size = 6, p = 0.3))

# Style the plot using color settings and ggplot2 functions:
lineup_histograms(tips, "total_bill",
                  dist = "gamma",
                  color_bars = "steelblue",
                  color_lines = "magenta") +
  ggplot2::theme_minimal()
```

lineup_qq

Check distributional assumptions using Q-Q plots and the lineup protocol.

Description

This function is used to quickly create lineup plots to check distributional assumptions using Q-Q plots. The null hypothesis is that the data follows the distribution specified by the `dist` argument. In the lineup protocol the plot of the real data is embedded amongst a field of plots of data generated to be consistent with some null hypothesis. If the observer can pick the real data as different from the others, this lends weight to the statistical significance of the structure in the plot. The protocol is described in Buja et al. (2009).

Usage

```
lineup_qq(
  data,
  variable,
  dist = NULL,
  params = NULL,
  color_points = "black",
```

```

    color_lines = "brown3",
    alpha_points = 0.5
  )

```

Arguments

data	a data frame.
variable	the name of the variable that should be plotted.
dist	the null distribution name. One of: "beta", "cauchy", "chi-squared", "exponential", "f", "gamma", "geometric", "log-normal", "lognormal", "logistic", "negative binomial", "normal", "poisson", "t", "uniform", "weibull"
params	list of parameters of distribution. If NULL, will use <code>fitdistr</code> to estimate them if possible. For uniform and beta distributions, the parameters must be specified. See <code>?dunif</code> and <code>?dbeta</code> for parameter names.
color_points	the color used for points. Can be a name or a color HEX code.
color_lines	the color used for reference lines.
alpha_points	the alpha (opacity) used for points (between 0 and 1, where 1 is opaque).

Details

19 null datasets are plotted together the the true data (randomly positioned) If you pick the real data as being noticeably different, then you have formally established that it is different to with p-value 0.05.

Run the decrypt message printed in the R Console to see which plot represents the true data.

Value

a ggplot

References

Buja, Cook, Hofmann, Lawrence, Lee, Swayne, Wickham. (2009). Statistical inference for exploratory data analysis and model diagnostics, *Phil. Trans. R. Soc. A*, 367, 4361-4383.

See Also

`null_dist`

Examples

```

data(tips)
lineup_qq(tips, "total_bill", dist = "normal") # Normal distribution
lineup_qq(tips, "total_bill", dist = "gamma") # Gamma distribution

# Some distributions require that the parameters be specified:
tips$proportion_tips <- tips$tip/(tips$total_bill+tips$tip)
lineup_qq(tips, "size", dist = "beta", params = list(shape1 = 0.1, shape2 = 0.2))

```

```
# Style the plot using color settings and ggplot2 functions:
lineup_qq(tips, "total_bill",
          dist = "gamma",
          color_points = "chocolate",
          color_lines = "cyan",
          alpha_points = 0.25) +
  ggplot2::theme_minimal()
```

lineup_residuals

Compare residual plots of a fitted model to plots of null residuals.

Description

This function is used to quickly create lineup version of the residual plots created by `plot.lm` and `ggfortify::autoplot.lm`; see Details for descriptions of these plots. In the lineup protocol the plot of the real data is embedded amongst a field of plots of data generated to be consistent with some null hypothesis. If the observer can pick the real data as different from the others, this lends weight to the statistical significance of the structure in the plot. The protocol is described in Buja et al. (2009).

Usage

```
lineup_residuals(
  model,
  type = 1,
  method = "rotate",
  color_points = "black",
  color_trends = "blue",
  color_lines = "brown3",
  alpha_points = 0.5,
  ...
)
```

Arguments

<code>model</code>	a model object fitted using lm .
<code>type</code>	type of plot: 1 = residuals vs fitted, 2 = normal Q-Q, 3 = scale-location, 4 = residuals vs leverage.
<code>method</code>	method for generating null residuals. Built in methods 'rotate', 'perm', 'pboot' and 'boot' are defined by resid_rotate , resid_perm , resid_pboot and resid_boot respectively. 'pboot' is always used for plots of type 2.
<code>color_points</code>	the color used for points in the plot. Can be a name or a color HEX code.
<code>color_trends</code>	the color used for trend curves in the plot.
<code>color_lines</code>	the color used for reference lines in the plot.
<code>alpha_points</code>	the alpha (opacity) used for points in the plot (between 0 and 1, where 1 is opaque).
<code>...</code>	other arguments passed onto method.

Details

Four types of plots are available:

1. Residual vs fitted. Null hypothesis: variable is linear combination of predictors.
2. Normal Q-Q plot. Null hypothesis: errors are normal. Always uses `method = "pboot"` to generate residuals under the null hypothesis.
3. Scale-location. Null hypothesis: errors are homoscedastic.
4. Residuals vs leverage. Used to identify points with high residuals and high leverage, which are likely to have a strong influence on the model fit.

19 null datasets are plotted together with the true data (randomly positioned). If you pick the real data as being noticeably different, then you have formally established that it is different with a p-value of 0.05. Run the `decrypt` message printed in the R Console to see which plot represents the true data.

If the null hypothesis in the type 1 plot is violated, consider using a different model. If the null hypotheses in the type 2 or 3 plots are violated, consider using bootstrap p-values; see [Section 8.1.5](#) of Thulin (2024) for details and recommendations.

Value

a `ggplot`

References

Buja, Cook, Hofmann, Lawrence, Lee, Swayne, Wickham. (2009). Statistical inference for exploratory data analysis and model diagnostics, *Phil. Trans. R. Soc. A*, 367, 4361-4383.

Thulin, M. (2024) *Modern Statistics with R*. Boca Raton: CRC Press. ISBN 9781032512440. <https://www.modernstatisticswithr.com/>

See Also

`null_lm`

Examples

```
data(tips)
x <- lm(tip ~ total_bill, data = tips)
lineup_residuals(x, type = 1) # Residuals vs Fitted
lineup_residuals(x, type = 2, method = "pboot") # Normal Q-Q plot
lineup_residuals(x, type = 4) # Residuals vs Leverage

# Style the plot using color settings and ggplot2 functions:
lineup_residuals(x, type = 3,
                 color_points = "skyblue",
                 color_trends = "darkorange") +
  ggplot2::theme_minimal()
```

null_dist	<i>Generate null data with a specific distribution.</i>
-----------	---

Description

Null hypothesis: variable has specified distribution

Usage

```
null_dist(var, dist, params = NULL)
```

Arguments

var	variable name
dist	distribution name. One of: beta, cauchy, chisq, exp, f, gamma, geom, lnorm, logis, nbinom, binom, norm, pois, t, unif, weibull
params	list of parameters of distribution. If NULL, will use fitdistr to estimate them.

Value

a function that given data generates a null data set. For use with [lineup](#) or [rorschach](#)

See Also

[null_permute](#), [null_lm](#)

Examples

```
dframe <- data.frame(x = rnorm(150))
library(ggplot2)
# three histograms of normally distributed values
ggplot(
  data=rorschach(method=null_dist("x", "norm"), n = 3, true=dframe)
) +
  geom_histogram(aes(x=x, y=..density..), binwidth=0.25) +
  facet_grid(.~.sample) +
  geom_density(aes(x=x), colour="steelblue", size=1)

# uniform distributions are not as easy to recognize as such
dframe$x = runif(150)
ggplot(
  data=rorschach(method=null_dist("x", "uniform",
    params=list(min=0, max=1)),
    n = 3, true=dframe)) +
  geom_histogram(aes(x=x, y=..density..), binwidth=0.1) +
  facet_grid(.~.sample) +
  geom_density(aes(x=x), colour="steelblue", size=1)
```

null_lm	<i>Generate null data with null residuals from a model.</i>
---------	---

Description

Null hypothesis: variable is linear combination of predictors

Usage

```
null_lm(f, method = "rotate", additional = FALSE, ...)
```

Arguments

f	model specification formula, as defined by lm
method	method for generating null residuals. Built in methods 'rotate', 'perm', 'pboot' and 'boot' are defined by resid_rotate , resid_perm , resid_pboot and resid_boot respectively
additional	whether to compute additional measures: standardized residuals and leverage
...	other arguments passed onto method.

Value

a function that given data generates a null data set. For use with [lineup](#) or [rorschach](#)

See Also

[null_permute](#), [null_dist](#)

Examples

```
data(tips)
x <- lm(tip ~ total_bill, data = tips)
tips.reg <- data.frame(tips, .resid = residuals(x), .fitted = fitted(x))
library(ggplot2)
ggplot(lineup(null_lm(tip ~ total_bill, method = 'rotate'), tips.reg)) +
  geom_point(aes(x = total_bill, y = .resid)) +
  facet_wrap(~ .sample)
```

null_permute	<i>Generate null data by permuting a variable.</i>
--------------	--

Description

Null hypothesis: variable is independent of others

Usage

```
null_permute(var)
```

Arguments

var	name of variable to permute
-----	-----------------------------

Value

a function that given data generates a null data set. For use with [lineup](#) or [rorschach](#)

See Also

[null_lm](#), [null_dist](#)

Examples

```
data(mtcars)
library(ggplot2)
ggplot(data=rorschach(method=null_permute("mpg"), n = 3, true=mtcars)) +
  geom_boxplot(aes(x=factor(cyl), y=mpg, fill=factor(cyl))) +facet_grid(.~.sample) +
  theme(legend.position="none", aspect.ratio=1)
```

null_ts	<i>Generate null data by simulating from a time series model.</i>
---------	---

Description

Null hypothesis: data follows a time series model using auto.arima from the forecast package

Usage

```
null_ts(var, modelfn)
```

Arguments

var	variable to model as a time series
modelfn	method for simulating from ts model.

Value

a function that given data generates a null data set. For use with [lineup](#) or [rorschach](#)

See Also

[null_model](#)

Examples

```
require(forecast)
require(ggplot2)
require(dplyr)
data(aud)
l <- lineup(null_ts("rate", auto.arima), aud)
ggplot(l, aes(x=date, y=rate)) + geom_line() +
  facet_wrap(~.sample, scales="free_y") +
  theme(axis.text = element_blank()) +
  xlab("") + ylab("")
l_dif <- l %>%
  group_by(.sample) %>%
  mutate(d=c(NA,diff(rate))) %>%
  ggplot(aes(x=d)) + geom_density() +
  facet_wrap(~.sample)
```

opt_bin_diff

Finds the number of bins in x and y direction which gives the maximum binned distance.

Description

This function finds the optimal number of bins in both x and y direction which should be used to calculate the binned distance. The binned distance is calculated for each combination of provided choices of number of bins in x and y direction and finds the difference using `calc_diff` for each combination. The combination for which the difference is maximum should be used.

Usage

```
opt_bin_diff(
  lineup.dat,
  var,
  xlow,
  xhigh,
  ylow,
  yhigh,
  pos,
  plot = FALSE,
  m = 20
)
```

Arguments

lineup.dat	lineup data to get the lineup
var	a list of names of the variables to be used to calculate the difference
xlow	the lowest value of number of bins on the x-direction
xhigh	the highest value of number of bins on the x-direction
ylow	the lowest value of number of bins on the y-direction
yhigh	the highest value of number of bins on the y-direction
pos	position of the true plot in the lineup
plot	LOGICAL; if true, returns a tile plot for the combinations of number of bins with the differences as weights
m	number of plots in the lineup, by default $m = 20$

Value

a dataframe with the number of bins and differences the maximum mean distance of the null plots

Examples

```
if(require('dplyr')){
  opt_bin_diff(lineup(null_permute('mpg'), mtcars, pos = 1), var = c('mpg', 'wt'),
  2, 5, 4, 8, pos = 1, plot = TRUE, m = 8)
}
```

pvisual

P-value calculations.

Description

These set of functions allow the user to calculate a p-value from the lineup after it has been evaluated by K independent observers. The different functions accommodate different lineup construction and showing to observers. Details are in the papers Majumder et al (2012) JASA, and Hofmann et al (2015). We distinguish between three different scenarios:

- Scenario I: in each of K evaluations a different data set and a different set of $(m-1)$ null plots is shown.
- Scenario II: in each of K evaluations the same data set but a different set of $(m-1)$ null plots is shown.
- Scenario III: the same lineup, i.e. same data and same set of null plots, is shown to K different observers.

Usage

```
pvisual(
  x,
  K,
  m = 20,
  N = 10000,
  type = "scenario3",
  xp = 1,
  target = 1,
  upper.tail = TRUE
)
```

Arguments

x	number of observed picks of the data plot
K	number of evaluations
m	size of the lineup
N	MC parameter: number of replicates on which MC probabilities are based. Higher number of replicates will decrease MC variability.
type	type of simulation used: scenario 3 assumes that the same lineup is shown in all K evaluations
xp	exponent used, defaults to 1
target	integer value identifying the location of the data plot
upper.tail	compute probabilities $P(X \geq x)$. Be aware that the use of this parameter is not consistent with the other distribution functions in base. There, a value of $P(X > x)$ is computed for upper.tail=TRUE.

Value

Vector/data frame. For comparison a p value based on a binomial distribution is provided as well.

Examples

```
pvisual(15, 20, m=3) # triangle test
```

 reg_dist

Distance based on the regression parameters

Description

Dataset X is binned into 5 bins in x-direction. A regression line is fitted to the data in each bin and the regression coefficients are noted. Same is done for dataset PX. An euclidean distance is calculated between the two sets of regression parameters. If the relationship between X and PX looks linear, number of bins should be equal to 1.

Usage

```
reg_dist(X, PX, nbins = 1, intercept = TRUE, scale = TRUE)
```

Arguments

X	a data.frame with two variables, the first column giving the explanatory variable and the second column giving the response variable
PX	another data.frame with two variables, the first column giving the explanatory variable and the second column giving the response variable
nbins	number of bins on the x-direction, by default nbins = 1
intercept	include the distances between intercepts?
scale	logical value: should the variables be scaled before computing regression coefficients?

Value

distance between X and PX

Examples

```
with(mtcars, reg_dist(data.frame(wt, mpg), data.frame(sample(wt), mpg)))
```

resid_boot

Bootstrap residuals.

Description

For use with [null_lm](#)

Usage

```
resid_boot(model, data)
```

Arguments

model	to extract residuals from
data	used to fit model

resid_pboot	<i>Parametric bootstrap residuals.</i>
-------------	--

Description

For use with [null_lm](#)

Usage

```
resid_pboot(model, data)
```

Arguments

model	to extract residuals from
data	used to fit model

resid_perm	<i>Permutation residuals.</i>
------------	-------------------------------

Description

For use with [null_lm](#)

Usage

```
resid_perm(model, data)
```

Arguments

model	to extract residuals from
data	used to fit model

resid_rotate	<i>Rotation residuals.</i>
--------------	----------------------------

Description

For use with [null_lm](#)

Usage

```
resid_rotate(model, data)
```

Arguments

model	to extract residuals from
data	used to fit model

resid_sigma	<i>Residuals simulated by a normal model, with specified sigma</i>
-------------	--

Description

For use with [null_lm](#)

Usage

```
resid_sigma(model, data, sigma = 1)
```

Arguments

model	to extract residuals from
data	used to fit model
sigma	a specific sigma to model

rorschach	<i>The Rorschach protocol.</i>
-----------	--------------------------------

Description

This protocol is used to calibrate the eyes for variation due to sampling. All plots are typically null data sets, data that is consistent with a null hypothesis. The protocol is described in Buja, Cook, Hofmann, Lawrence, Lee, Swayne, Wickham (2009) Statistical inference for exploratory data analysis and model diagnostics, Phil. Trans. R. Soc. A, 367, 4361-4383.

Usage

```
rorschach(method, true = NULL, n = 20, p = 0)
```

Arguments

method	method for generating null data sets
true	true data set. If NULL, find_plot_data will attempt to extract it from the current ggplot2 plot.
n	total number of samples to generate (including true data)
p	probability of including true data with null data.

sample_size	<i>Sample size calculator</i>
-------------	-------------------------------

Description

This function calculates a table of sample sizes for with an experiment, given a lineup size, and estimates of the detection rate.

Usage

```
sample_size(n = 53:64, m = 20, pA = seq(1/20, 1/3, 0.01), conf = 0.95)
```

Arguments

n	range of sample sizes to check, default is 53:64
m	linup size, default 20
pA	range of estimated detection rates to consider, default is seq(1/20, 1/3, 0.01)
conf	confidence level to use to simulate from binomial

Examples

```
pow <- sample_size()
pow
library(ggplot2)
library(viridis)
ggplot(pow, aes(x=n, y=pA, fill=prob, group=pA)) +
  geom_tile() +
  scale_fill_viridis_c("power") +
  ylab("detect rate (pA)") + xlab("sample size (n)") +
  theme_bw()
```

sep_dist	<i>Distance based on separation of clusters</i>
----------	---

Description

The separation between clusters is defined by the minimum distances of a point in the cluster to a point in another cluster. The number of clusters are provided. If not, the hierarchical clustering method is used to obtain the clusters. The separation between the clusters for dataset X is calculated. Same is done for dataset PX. An euclidean distance is then calculated between these separation for X and PX.

Usage

```
sep_dist(X, PX, clustering = FALSE, nclust = 3, type = "separation")
```

Arguments

X	a data.frame with two or three columns, the first two columns providing the dataset
PX	a data.frame with two or three columns, the first two columns providing the dataset
clustering	LOGICAL; if TRUE, the third column is used as the clustering variable, by default FALSE
nclust	the number of clusters to be obtained by hierarchical clustering, by default nclust = 3
type	character string to specify which measure to use for distance, see ?cluster.stats for details

Value

distance between X and PX

Examples

```
if(require('fpc')) {
  with(mtcars, sep_dist(data.frame(wt, mpg, as.numeric(as.factor(mtcars$cyl))),
                        data.frame(sample(wt), mpg, as.numeric(as.factor(mtcars$cyl))),
                        clustering = TRUE))
}

if (require('fpc')) {
  with(mtcars, sep_dist(data.frame(wt, mpg, as.numeric(as.factor(mtcars$cyl))),
                        data.frame(sample(wt), mpg, as.numeric(as.factor(mtcars$cyl))),
                        nclust = 3))
}
```

theme_strip

A theme to minimally strip away the context

Description

Note this is not a complete theme hence why there are no arguments.

Usage

```
theme_strip()
```

Examples

```
library(ggplot2)
ggplot(cars, aes(dist, speed)) + theme_strip()
```

tips

Tipping data

Description

One waiter recorded information about each tip he received over a period of a few months working in one restaurant. He collected several variables:

Usage

tips

Format

A data frame with 244 rows and 7 variables

Details

- tip in dollars,
- bill in dollars,
- sex of the bill payer,
- whether there were smokers in the party,
- day of the week,
- time of day,
- size of the party.

In all he recorded 244 tips. The data was reported in a collection of case studies for business statistics (Bryant & Smith 1995).

References

Bryant, P. G. and Smith, M (1995) *Practical Data Analysis: Case Studies in Business Statistics*. Homewood, IL: Richard D. Irwin Publishing:

turk_results

Sample turk results

Description

Subset of data from a Turk experiment, used to show how to compute power of a lineup

uni_dist	<i>Distance for univariate data</i>
----------	-------------------------------------

Description

The first four moments is calculated for data X and data PX. An euclidean distance is calculated between these moments for X and PX.

Usage

```
uni_dist(X, PX)
```

Arguments

X	a data.frame where the first column is only used
PX	another data.frame where the first column is only used

Value

distance between X and PX

Examples

```
if(require('moments')){uni_dist(rnorm(100), rpois(100, 2))}
```

visual_power	<i>Power calculations.</i>
--------------	----------------------------

Description

This function simply counts the proportion of people who selected the data plot, in a set of lineups. It adjusts for multiple picks by the same individual, by weighting by the total number of choices.

Usage

```
visual_power(data, m = 20)
```

Arguments

data	summary of the results, containing columns id, pic_id, response, detected
m	size of the lineup

Value

vector of powers for each pic_id

Examples

```
data(turk_results)
visual_power(turk_results)
```

wasps

Wasp gene expression data.

Description

Data from Toth et al (2010) used in Niladri Roy et al (2015)

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