

Package ‘mbRes’

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Type Package

Title Exploration of Multiple Biomarker Responses using Effect Size

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Suggests RProbSup (>= 3.0)

BugReports <https://github.com/phamdn/mbRes/issues>

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mbRes-package	<i>mbRes: Exploration of Multiple Biomarker Responses using Effect Size</i>
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Description

Summarize multiple biomarker responses of aquatic organisms to contaminants using Cliff's delta, as described in Pham & Sokolova (2023) [doi:10.1002/team.4676](#).

Guidelines

`mbr` and `visual` are the main functions to compute and visualize Cliff's delta and S-value which are results of `cliff` and `resampling`. `setpop`, `simul`, and `plotsam` simulate and visualize a hypothetical dataset. `compare` compares the results of Cliff's delta and two other integrated indices published earlier (i.e., RSI and IBR, see `blaise2002` and `beliaeff2002`). The others (`ggheat` and `ggdot`) are helper functions and are not meant to be called directly by users.

Updates

`mbr.cliff` and `mbr.glass` simply compute and visualize Cliff's delta and Glass's delta.

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beliaeoff2002

Compute Integrated Biomarker Index

Description

beliaeoff2002 calculates IBR in the hypothetical case study. This is not meant to be called directly.

Usage

`beliaeoff2002(sam_mean)`

Arguments

`sam_mean` a data frame, the third output of `simul`.

Value

beliaeoff2002 returns a data frame of IBR.

References

Beliaeoff, B., & Burgeot, T. (2002). Integrated biomarker response: A useful tool for ecological risk assessment. Environmental Toxicology and Chemistry, 21(6), 1316–1322. [doi:10.1002/etc.5620210629](https://doi.org/10.1002/etc.5620210629).

blaise2002

Compute Rank Sum Biomarker Index

Description

blaise2002 calculates RSI in the hypothetical case study. This is not meant to be called directly.

Usage

`blaise2002(sam, sam_mean)`

Arguments

`sam` a data frame, the first output of `simul`.

`sam_mean` a data frame, the third output of `simul`.

Value

`blaise2002` returns a data frame of RSI.

References

Blaise, C., Gagné, F., Pellerin, J., Hansen, P.-D., & Trottier, S. (2002). Molluscan shellfish biomarker study of the Quebec, Canada, Saguenay Fjord with the soft-shell clam, *Mya arenaria*. *Environmental Toxicology*, 17(3), 170–186. doi:[10.1002/tox.10048](https://doi.org/10.1002/tox.10048).

`cliff`

Compute Effect Size

Description

`cliff` calculates Cliff's delta statistic using the rank sum method.

Usage

`cliff(v1, v0)`

Arguments

<code>v1</code>	a vector, biomarker values from the treatment group.
<code>v0</code>	a vector, biomarker values from the control group.

Value

`cliff` returns a numeric that is the Cliff's delta of the treatment group.

References

Cliff, N. (1993). Dominance statistics: Ordinal analyses to answer ordinal questions. *Psychological Bulletin*, 114(3), 494–509. doi:[10.1037/00332909.114.3.494](https://doi.org/10.1037/00332909.114.3.494).

Vargha, A., & Delaney, H. D. (2000). A Critique and Improvement of the CL Common Language Effect Size Statistics of McGraw and Wong. *Journal of Educational and Behavioral Statistics*, 25(2), 101–132. doi:[10.3102/10769986025002101](https://doi.org/10.3102/10769986025002101).

Ruscio, J., & Mullen, T. (2012). Confidence Intervals for the Probability of Superiority Effect Size Measure and the Area Under a Receiver Operating Characteristic Curve. *Multivariate Behavioral Research*, 47(2), 201–223. doi:[10.1080/00273171.2012.658329](https://doi.org/10.1080/00273171.2012.658329).

See Also

[CalcA1](#).

Examples

```
set.seed(1)
setting <- setpop()
temp <- simul(setting$pop_mean)
cliff(subset(temp$sam, Site == "S1", Bmk1, drop = TRUE),
subset(temp$sam, Site == "S0", Bmk1, drop = TRUE))
```

compare

Compare RSI, IBR, and Cliff's delta

Description

compare calculates RSI assigned values, IBR translated scores, and Cliff's delta in the hypothetical case study.

Usage

```
compare(sam, sam_mean)
```

Arguments

sam	a data frame, the first output of simul .
sam_mean	a data frame, the third output of simul .

Value

compare returns a list of length 5:

blaise	RSI assigned values and final RSI.
beliaeaff	IBR translated scores and final IBR.
pham	Cliff's delta and the average of absolute Cliff's delta.
fig1	ggplot object of comparisions among RSI assigned values, IBR translated scores, and Cliff's delta.
fig2	ggplot object of comparision among RSI, IBR, and the average of absolute Cliff's delta.

References

Blaise, C., Gagné, F., Pellerin, J., Hansen, P.-D., & Trottier, S. (2002). Molluscan shellfish biomarker study of the Quebec, Canada, Saguenay Fjord with the soft-shell clam, Mya arenaria. Environmental Toxicology, 17(3), 170–186. [doi:10.1002/tox.10048](https://doi.org/10.1002/tox.10048).

Beliaeaff, B., & Burgeot, T. (2002). Integrated biomarker response: A useful tool for ecological risk assessment. Environmental Toxicology and Chemistry, 21(6), 1316–1322. [doi:10.1002/etc.5620210629](https://doi.org/10.1002/etc.5620210629).

Examples

```
set.seed(1)
setting <- setpop()
temp <- simul(setting$pop_mean)
compare(temp$sam, temp$sam_mean)
#might take more than 5s in some machines
```

ggdot

Make Dot Plot

Description

ggdot creates dot plot of the average of absolute Cliff's delta. This is not meant to be called directly.

Usage

```
ggdot(dat, hax, vax)
```

Arguments

dat	a data frame with at least two columns.
hax	a character, name of the column to be used as the horizontal axis.
vax	a character, name of the column to be used as the vertical axis.

Value

ggdot returns a ggplot object.

ggheat

Make Heatmap

Description

ggheat creates heatmaps of the Cliff's delta and S-value. This is not meant to be called directly.

Usage

```
ggheat(
  dat,
  hax,
  vax,
  cell,
  nm,
  lim,
  lo,
  hi,
  diverging = FALSE,
  env = parent.frame()
)
```

Arguments

dat	a data frame with at least three columns.
hax	a character, name of the column to be used as the horizontal axis.
vax	a character, name of the column to be used as the vertical axis.
cell	a character, name of the column to be used as the cells.
nm	a character, name of the heatmap.
lim	a numeric vector, limits of the color scale.
lo	a character, color of the color scale low end.
hi	a character, color of the color scale high end.
diverging	a logical, whether to use diverging color gradient.
env	an environment, to access outer scope variables.

Value

ggheat returns a ggplot object.

mbr

Compute Cliff's delta and S-value
Description

mbr summarizes Cliff's delta and S-value for multiple groups and multiple biomarkers.

Usage

```
mbr(df)
```

Arguments

- df** a data frame with the name of experimental groups or biomonitoring sites as the first column and the measurement of biomarkers as the remaining columns.

Details

The header of the first column can be any character, for example, 'group' or 'site'. The first name appearing in the first column will determine the control group or the reference site. The other names will be treatment groups or test sites. The header of the remaining columns will define the list of biomarkers.

Value

mbr returns a list of length 3:

- | | |
|-------------|--|
| mess | a list of length 3 confirms the information about df. |
| es | a data frame with 9 columns:

test_site treatment groups or test sites.
ref_site control group or reference site.
t_size the sample size of treatment group or test sites.
r_size the sample size of control group or reference site.
biomarker individual biomarker.
delta the Cliff's delta of treatment group or reference site.
delta.abs the absolute Cliff's delta.
pval the P-Value.
sval the surprisal or S-Value. |
| idx | a data frame summarizes delta.abs and their average. |

Examples

```
set.seed(1)
setting <- setpop()
temp <- simul(setting$pop_mean)
mbr(temp$sam)
#might take more than 5s in some machines
```

mbr.cliff*Compute Cliff's delta simplified*

Description

`mbr.cliff` summarizes Cliff's delta for multiple groups and multiple biomarkers.

Usage

```
mbr.cliff(df)
```

Arguments

`df` a data frame with the name of experimental groups or biomonitoring sites as the first column and the measurement of biomarkers as the remaining columns.

Examples

```
set.seed(1)
setting <- setpop()
temp <- simul(setting$pop_mean)
mbr.cliff(temp$sam)
#might take more than 5s in some machines
```

mbr.glass*Compute Glass's delta simplified*

Description

`mbr.glass` summarizes Glass's delta for multiple groups and multiple biomarkers.

Usage

```
mbr.glass(df)
```

Arguments

`df` a data frame with the name of experimental groups or biomonitoring sites as the first column and the measurement of biomarkers as the remaining columns.

Examples

```
set.seed(1)
setting <- setpop()
temp <- simul(setting$pop_mean)
mbr.glass(temp$sam)
#might take more than 5s in some machines
```

plotsam

Visualize Hypothetical Samples

Description

`plotsam` plots the sample dataset of biomarker responses. This is used for the hypothetical case study.

Usage

```
plotsam(pop_mean_long, pop_profile, sam_long)
```

Arguments

<code>pop_mean_long</code>	a data frame, the second output of <code>setpop</code> .
<code>pop_profile</code>	a data frame, the third output of <code>setpop</code> .
<code>sam_long</code>	a data frame, the second output of <code>simul</code> .

Value

`plotsam` returns a ggplot object.

Examples

```
set.seed(1)
setting <- setpop()
temp <- simul(setting$pop_mean)
plotsam(setting$pop_mean_long, setting$pop_profile, temp$sam_long)
```

resampling

Measure Statistical Uncertainty

Description

resampling performs randomization test to calculate P-value and S-value.

Usage

```
resampling(v1, v0, nrand = 1999, seed = 1)
```

Arguments

v1	a vector, biomarker values from the treatment group.
v0	a vector, biomarker values from the control group.
nrand	an integer, the number of randomization samples. The default value is 1999.
seed	an integer, the seed for random number generation. Setting a seed ensures the reproducibility of the result. See set.seed for more details.

Value

resampling returns a one-row data frame with 3 numerics:

delta	the Cliff's delta of the treatment group.
pval	the observed P-value p under the null hypothesis.
sval	the S-value s calculated from P-value p.

References

Greenland, S. (2019). Valid P-Values Behave Exactly as They Should: Some Misleading Criticisms of P-Values and Their Resolution With S-Values. *The American Statistician*, 73(sup1), 106–114. doi:[10.1080/00031305.2018.1529625](https://doi.org/10.1080/00031305.2018.1529625).

Phipson, B., & Smyth, G. K. (2010). Permutation P-values Should Never Be Zero: Calculating Exact P-values When Permutations Are Randomly Drawn. *Statistical Applications in Genetics and Molecular Biology*, 9(1). doi:[10.2202/15446115.1585](https://doi.org/10.2202/15446115.1585).

See Also

[A1](#).

Examples

```
set.seed(1)
setting <- setpop()
temp <- simul(setting$pop_mean)
resampling(subset(temp$sam, Site == "S1", Bmk1, drop = TRUE),
subset(temp$sam, Site == "S0", Bmk1, drop = TRUE))
```

setpop

Define Hypothetical Populations

Description

`setpop` sets the true means of biomarker responses in populations. This is used for the hypothetical case study.

Usage

```
setpop()
```

Value

`setpop` returns a list of length 3:

<code>pop_mean</code>	true means of biomarker responses in populations.
<code>pop_mean_long</code>	true means of biomarker responses in long format.
<code>pop_profile</code>	profile of biomarkers.

simul

Generate Hypothetical Samples

Description

`simul` yields a sample dataset of biomarker responses. This is used for the hypothetical case study.

Usage

```
simul(pop_mean, size = 75)
```

Arguments

<code>pop_mean</code>	a data frame, the first output of <code>setpop</code> .
<code>size</code>	an integer, the sample size.

Value

`simul` returns a list of length 3:

<code>sam</code>	sample dataset.
<code>sam_long</code>	sample dataset in long format.
<code>sam_mean</code>	sample means of biomarker responses.

<code>visual</code>	<i>Visualize Cliff's delta and S-value</i>
---------------------	--

Description

`visual` plots Cliff's delta and S-value for multiple groups and multiple biomarkers.

Usage

```
visual(rs, rotate = FALSE, display = TRUE)
```

Arguments

<code>rs</code>	a list, output of <code>mbr</code> .
<code>rotate</code>	a logical, whether to rotate the biomarker labels in figures.
<code>display</code>	a logical, whether to display cell values in heatmaps.

Value

`visual` returns a list of ggplot objects:

<code>fig.delta</code>	heatmap of Cliff's delta.
<code>fig.sval</code>	heatmap of S-value.
<code>fig.avg</code>	dot plot of the average of absolute Cliff's delta.
<code>mbr_fig</code>	combined heatmaps of Cliff's delta and S-value.

Examples

```
set.seed(1)
setting <- setpop()
temp <- simul(setting$pop_mean)
mbr_result <- mbr(temp$sam)
visual(mbr_result)
#might take more than 5s in some machines
```

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