Package ‘riskclustr’

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Type Package

Title Functions to Study Etiologic Heterogeneity

Version 0.4.1

Description A collection of functions related to the study of etiologic heterogeneity both across disease subtypes and across individual disease markers. The included functions allow one to quantify the extent of etiologic heterogeneity in the context of a case-control study, and provide p-values to test for etiologic heterogeneity across individual risk factors. Begg CB, Zabor EC, Bernstein JL, Bernstein L, Press MF, Seshan VE (2013) <doi:10.1002/sim.5902>.

Depends R (>= 4.0)

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BugReports https://github.com/zabore/riskclustr/issues

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| d | Estimate the incremental explained risk variation in a case-control study |
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**Description**

d estimates the incremental explained risk variation across a set of pre-specified disease subtypes in a case-control study. This function takes the name of the disease subtype variable, the number of disease subtypes, a list of risk factors, and a wide dataset, and does the needed transformation on the dataset to get the correct format. Then the polytomous logistic regression model is fit using `mlogit`, and D is calculated based on the resulting risk predictions.

**Usage**

d(label, M, factors, data)

**Arguments**

- **label**: the name of the subtype variable in the data. This should be a numeric variable with values 0 through M, where 0 indicates control subjects. Must be supplied in quotes, e.g. `label = "subtype"`. quotes.
- **M**: is the number of subtypes. For M>=2.
- **factors**: a list of the names of the binary or continuous risk factors. For binary risk factors the lowest level will be used as the reference level. e.g. `factors = list("age", "sex", "race")`.
- **data**: the name of the dataframe that contains the relevant variables.

**References**

**Examples**

```r
d(d(
    label = "subtype",
    M = 4,
    factors = list("x1", "x2", "x3"),
    data = subtype_data
)
)
```

**dstar**

*Estimate the incremental explained risk variation in a case-only study*

**Description**

*dstar* estimates the incremental explained risk variation across a set of pre-specified disease subtypes in a case-only study. The highest frequency level of label is used as the reference level, for stability. This function takes the name of the disease subtype variable, the number of disease subtypes, a list of risk factors, and a wide case-only dataset, and does the needed transformation on the dataset to get the correct format. Then the polytomous logistic regression model is fit using `mlogit`, and $D^*$ is calculated based on the resulting risk predictions.

**Usage**

```r
dstar(label, M, factors, data)
```

**Arguments**

- **label**
  
  the name of the subtype variable in the data. This should be a numeric variable with values 0 through M, where 0 indicates control subjects. Must be supplied in quotes, e.g. `label = "subtype"`. quotes.

- **M**
  
  is the number of subtypes. For M=2.

- **factors**
  
  a list of the names of the binary or continuous risk factors. For binary risk factors the lowest level will be used as the reference level. e.g. `factors = list("age", "sex", "race")`.

- **data**
  
  the name of the case-only dataframe that contains the relevant variables.

**References**

Examples

```r
# Exclude controls from data as this is a case-only calculation
dstar(
    label = "subtype",
    M = 4,
    factors = list("x1", "x2", "x3"),
    data = subtype_data[subtype_data$subtype > 0, ]
)
```

dh_test_marker

**Test for etiologic heterogeneity of risk factors according to individual
disease markers in a case-control study**

Description

dh_test_marker takes a list of individual disease markers, a list of risk factors, a variable name
denoting case versus control status, and a dataframe, and returns results related to the question of
whether each risk factor differs across levels of the disease subtypes and the question of whether
each risk factor differs across levels of each individual disease marker of which the disease subtypes
are comprised. Input is a dataframe that contains the individual disease markers, the risk factors of
interest, and an indicator of case or control status. The disease markers must be binary and must
have levels 0 or 1 for cases. The disease markers should be left missing for control subjects. For
categorical disease markers, a reference level should be selected and then indicator variables for
each remaining level of the disease marker should be created. Risk factors can be either binary
or continuous. For categorical risk factors, a reference level should be selected and then indicator
variables for each remaining level of the risk factor should be created.

Usage

dh_test_marker(markers, factors, case, data, digits = 2)

Arguments

- **markers**: a list of the names of the binary disease markers. Each must have levels 0 or
  1 for case subjects. This value will be missing for all control subjects. e.g.
  `markers = list("marker1", "marker2")`

- **factors**: a list of the names of the binary or continuous risk factors. For binary risk factors
  the lowest level will be used as the reference level. e.g. `factors = list("age", "sex", "race")`

- **case**: denotes the variable that contains each subject’s status as a case or control. This
  value should be 1 for cases and 0 for controls. Argument must be supplied in
  quotes, e.g. `case = "status"`. 

- **data**: the name of the dataframe that contains the relevant variables.

- **digits**: the number of digits to round the odds ratios and associated confidence intervals,
  and the estimates and associated standard errors. Defaults to 2.
Value

Returns a list.

beta is a matrix containing the raw estimates from the polytomous logistic regression model fit with mlogit with a row for each risk factor and a column for each disease subtype.

beta_se is a matrix containing the raw standard errors from the polytomous logistic regression model fit with mlogit with a row for each risk factor and a column for each disease subtype.

eh_pval is a vector of unformatted p-values for testing whether each risk factor differs across the levels of the disease subtype.

gamma is a matrix containing the estimated disease marker parameters, obtained as linear combinations of the beta estimates, with a row for each risk factor and a column for each disease marker.

gamma_se is a matrix containing the estimated disease marker standard errors, obtained based on a transformation of the beta standard errors, with a row for each risk factor and a column for each disease marker.

gamma_p is a matrix of p-values for testing whether each risk factor differs across levels of each disease marker, with a row for each risk factor and a column for each disease marker.

or_ci_p is a dataframe with the odds ratio (95%) factor/subtype combination, as well as a column of formatted etiologic heterogeneity p-values.

beta_se_p is a dataframe with the estimates (SE) for each risk factor/subtype combination, as well as a column of formatted etiologic heterogeneity p-values.

gamma_se_p is a dataframe with disease marker estimates (SE) and their associated p-values.

Author(s)

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Examples

# Run for two binary tumor markers, which will combine to form four subtypes
eh_test_marker(
    markers = list("marker1", "marker2"),
    factors = list("x1", "x2", "x3"),
    case = "case",
    data = subtype_data,
    digits = 2
)

---

**eh_test_subtype**

*Test for etiologic heterogeneity of risk factors according to disease subtypes in a case-control study*
Description

`eh_test_subtype` takes the name of the variable containing the pre-specified subtype labels, the number of subtypes, a list of risk factors, and the name of the dataframe and returns results related to the question of whether each risk factor differs across levels of the disease subtypes. Input is a dataframe that contains the risk factors of interest and a variable containing numeric class labels that is 0 for control subjects. Risk factors can be either binary or continuous. For categorical risk factors, a reference level should be selected and then indicator variables for each remaining level of the risk factor should be created. Categorical risk factors entered as is will be treated as ordinal. The multinomial logistic regression model is fit using `mlogit`.

Usage

```r
eh_test_subtype(label, M, factors, data, digits = 2)
```

Arguments

- `label`: the name of the subtype variable in the data. This should be a numeric variable with values 0 through M, where 0 indicates control subjects. Must be supplied in quotes, e.g. `label = "subtype"`.
- `M`: is the number of subtypes. For M>=2.
- `factors`: a list of the names of the binary or continuous risk factors. For binary or categorical risk factors the lowest level will be used as the reference level. e.g. `factors = list("age", "sex", "race")`.
- `data`: the name of the dataframe that contains the relevant variables.
- `digits`: the number of digits to round the odds ratios and associated confidence intervals, and the estimates and associated standard errors. Defaults to 2.

Value

Returns a list.

- `beta`: is a matrix containing the raw estimates from the polytomous logistic regression model fit with `mlogit` with a row for each risk factor and a column for each disease subtype.
- `beta_se`: is a matrix containing the raw standard errors from the polytomous logistic regression model fit with `mlogit` with a row for each risk factor and a column for each disease subtype.
- `eh_pval`: is a vector of unformatted p-values for testing whether each risk factor differs across the levels of the disease subtype.
- `or_ci_p`: is a dataframe with the odds ratio (95\% factor/subtype combination, as well as a column of formatted etiologic heterogeneity p-values.
- `beta_se_p`: is a dataframe with the estimates (SE) for each risk factor/subtype combination, as well as a column of formatted etiologic heterogeneity p-values.
- `var_covar`: contains the variance-covariance matrix associated with the model estimates contained in `beta`.

Author(s)

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optimal_kmeans_d

Examples

```r
eh_test_subtype(
    label = "subtype",
    M = 4,
    factors = list("x1", "x2", "x3"),
    data = subtype_data,
    digits = 2
)
```

optimal_kmeans_d Obtain optimal D solution based on k-means clustering of disease marker data in a case-control study

Description

optimal_kmeans_d applies k-means clustering using the kmeans function with many random starts. The D value is then calculated for the cluster solution at each random start using the d function, and the cluster solution that maximizes D is returned, along with the corresponding value of D. In this way the optimally etiologically heterogeneous subtype solution can be identified from possibly high-dimensional disease marker data.

Usage

```r
optimal_kmeans_d(markers, M, factors, case, data, nstart = 100, seed = NULL)
```

Arguments

- **markers**: a vector of the names of the disease markers. These markers should be of a type that is suitable for use with kmeans clustering. All markers will be missing for control subjects. e.g. `markers = c("marker1", "marker2")`
- **M**: is the number of clusters to identify using kmeans clustering. For M>=2.
- **factors**: a list of the names of the binary or continuous risk factors. For binary risk factors the lowest level will be used as the reference level. e.g. `factors = list("age", "sex", "race")`
- **case**: denotes the variable that contains each subject’s status as a case or control. This value should be 1 for cases and 0 for controls. Argument must be supplied in quotes, e.g. `case = "status"`.
- **data**: the name of the dataframe that contains the relevant variables.
- **nstart**: the number of random starts to use with kmeans clustering. Defaults to 100.
- **seed**: an integer argument passed to set.seed. Default is NULL. Recommended to set in order to obtain reproducible results.
Value

Returns a list

- `optimal_d` The D value for the optimal D solution
- `optimal_d_data` The original data frame supplied through the `data` argument, with a column called `optimal_d_label` added for the optimal D subtype label. This has the subtype assignment for cases, and is 0 for all controls.

References


Examples

```r
# Cluster 30 disease markers to identify the optimally
# etiologically heterogeneous 3-subtype solution
res <- optimal_kmeans_d(
  markers = c(paste0("y", seq(1:30))),
  M = 3,
  factors = list("x1", "x2", "x3"),
  case = "case",
  data = subtype_data,
  nstart = 100,
  seed = 81110224
)

# Look at the value of D for the optimal D solution
res["optimal_d"]

# Look at a table of the optimal D solution
table(res["optimal_d_data"]$optimal_d_label)
```

---

**posthoc_factor_test**

Post-hoc test to obtain overall p-value for a factor variable used in a `eh_test_subtype` fit.

**Description**

`posthoc_factor_test` takes a `eh_test_subtype` fit and returns an overall p-value for a specified factor variable.

**Usage**

`posthoc_factor_test(fit, factor, nlevels)`
**Arguments**

- **fit**: the resulting `eh_test_subtype` fit.
- **factor**: is the name of the factor variable of interest, supplied in quotes, e.g. `factor = "race"`. Only supports a single factor.
- **nlevels**: is the number of levels the factor variable in `factor` has.

**Value**

Returns a list.
- `pval` is a formatted p-value.
- `pval_raw` is the raw, unformatted p-value.

**Author(s)**

Emily C Zabor <zabore@mskcc.org>

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**subtype_data**

Simulated subtype data

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**Description**

A dataset containing 2000 patients: 1200 cases and 800 controls. There are four subtypes, and both numeric and character subtype labels. The subtypes are formed by cross-classification of two binary disease markers, disease marker 1 and disease marker 2. There are three risk factors, two continuous and one binary. One of the continuous risk factors and the binary risk factor are related to the disease subtypes. There are also 30 continuous tumor markers, 20 of which are related to the subtypes and 10 of which represent noise, which could be used in a clustering analysis.

**Usage**

`subtype_data`

**Format**

A data frame with 2000 rows—one row per patient

- **case**: Indicator of case control status, 1 for cases and 0 for controls
- **subtype**: Numeric subtype label, 0 for control subjects
- **subtype_name**: Character subtype label
- **marker1**: Disease marker 1
- **marker2**: Disease marker 2
- **x1**: Continuous risk factor 1
- **x2**: Continuous risk factor 2
- **x3**: Binary risk factor
y1 Continuous tumor marker 1
y2 Continuous tumor marker 2
y3 Continuous tumor marker 3
y4 Continuous tumor marker 4
y5 Continuous tumor marker 5
y6 Continuous tumor marker 6
y7 Continuous tumor marker 7
y8 Continuous tumor marker 8
y9 Continuous tumor marker 9
y10 Continuous tumor marker 10
y11 Continuous tumor marker 11
y12 Continuous tumor marker 12
y13 Continuous tumor marker 13
y14 Continuous tumor marker 14
y15 Continuous tumor marker 15
y16 Continuous tumor marker 16
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y26 Continuous tumor marker 26
y27 Continuous tumor marker 27
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