Package 'eat'

February 20, 2012

Type Package

Title eat

| Version 1.4.7 | | |
|---|--|--|
| Depends R(>= 2.14.0), sendmailR, gdata, xlsx, car, reshape, foreign,date, plyr, parallel | | |
| Date 2012-02-20 | | |
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| Description The package eat is designed to simplify data preparation and IRT modeling with the software ConQuest within the R programming environment. It includes routines for automation of data preprocessing and an interface to specify and run several IRT models. | | |
| License GPL | | |
| LazyLoad yes | | |
| LazyData yes | | |
| R topics documented: | | |
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Description

More about what it does (maybe more than one line) ~~ A concise (1-5 lines) description of the package ~~

Details

Package: eat Type: Version: Package

Date:

License: What license is it under?

LazyLoad: yes aggregateData 3

~~ An overview of how to use the package, including the most important functions ~~

Author(s)

Who wrote it

Maintainer: Who to complain to <yourfault@somewhere.net> ~~ The author and/or maintainer of the package ~~

References

~~ Literature or other references for background information ~~

aggregateData

Aggregate Datasets with Missing Values

Description

Aggregates datasets with constraints on missing values

Usage

aggregateData(dat, subunits, units, aggregatemissings = "use.default", rename = FALSE, recodedData

Arguments

dat A data frame.

subunits A data frame with subunit information. See 'Details'.

A data frame with unit information. See 'Details'.

aggregatemissings

Either the character string "use.default" or a $n \times n$ matrix with information

on how missing values should be aggregated. See 'Details'.

rename Should units with only one subunit be renamed to their unit name? Default is

FALSE.

recodedData Logical indicating whether colnames in dataset to aggregate are the subunit

names (as in subunits\$subunit) or recoded subunit names (as in subunits\$subunitRecoded).

Default is TRUE, meaning that colnames are recoded subitem names.

Details

aggregateData aggregates units in data frames with special consideration of missing values. The aggregation of missing values is specified in argument aggregatemissings.

The results of aggregateData will be written to a protocol file with sunk.

Examples of data frames subunits and units can be found via data(inputList).

Value

A data frame with aggregated units and, if rename = TRUE, renamed subunits.

4 asNumericIfPossible

Warning

Missings are only correctly aggregated if their values correspond to the values given in aggregatemissings. aggregateData does not check for value types or whether codes are valid. Use of checkData and recodeData before using aggregateData is therefore strongly recommended.

Author(s)

Nicole Haag, Anna Lenski

References

For missing types see http://code.google.com/p/zkdlib/wiki/MissingHandling

See Also

recodeData, checkData

Examples

```
data(inputDat)
data(inputList)

dat1 <- inputDat[[1]] # get first dataset from inputDat
datRec <- recodeData(dat1, inputList$values, inputList$subunits) # recode Data first
datAggr <- aggregateData(datRec, inputList$subunits, inputList$units, rename = TRUE, recodedData = TRUE)</pre>
```

 $as {\tt Numeric If Possible}$

Transform columns of a data.frame into numeric values if possible

Description

In contrast to as.numeric, Function transforms only "transformable" columns of a data.frame into numeric values (i.e. without creating NA when transformation fails. Non-transformable columns are maintained. Optionally, only a logical vector is given, indicating which columns are transformable.

Usage

```
asNumericIfPossible ( dataFrame, set.numeric = TRUE, transform.factors = FALSE, maintain.factor.
```

Arguments

dataFrame

A data.frame which columns should be transformed.

set.numeric

Logical: If TRUE, data.frame with transformed columns is returned. If FALSE, a

logical vector is returned, indicating which columns are transformable.

transform.factors

Logical: Should columns of class factor transformed? If FALSE, columns of class factor are maintained. If TRUE, columns of class factor are attempted to transform.

maintain.factor.scores

Logical. Only relevant if transform.factors = TRUE. If TRUE, the nominal values of the factor are transformed if possible. If FALSE, the integer numbers representing the factors' nominal values are returned. See details.

verbose

Logical: If TRUE, informations about the class of the columns in the data.frame are printed to the console.

Details

In R, factors may represent ordered categories or nominal variables. Depending on the meaning of the variable, a transformation of the nominal values (of a factor variable) to numeric values may be desirable or not. The arguments transform.factors and maintain.factor.scores serve to specify if and how factor variables should be transformed. See examples.

Value

Either a logic vector, indicating which columns in the data.frame are transformable according to the specified conditions, ora data.frame in which transformable columns are transformed.

Author(s)

Sebastian Weirich

Examples

```
( dat <- data.frame( X1 = c("1",NA,"0"), X2 = c("a",NA,"b"), X3 = c(TRUE,FALSE,FALSE), X4 = as.factor(
str(dat)
asNumericIfPossible(dat)
asNumericIfPossible(dat, transform.factors=TRUE, maintain.factor.scores=FALSE)
asNumericIfPossible(dat, transform.factors=TRUE, maintain.factor.scores=TRUE)</pre>
```

 $automate {\tt Conquest Model} \quad automate {\tt Conquest Model}$

Description

Wrapper function which calls several functions to build and write Conquest input files.

Usage

```
automateConquestModel(dataset, ID, regression=NULL, DIF=NULL, group.var=NULL, weight=NULL, testitems, na=list(items=NULL, DIF=NULL, HG=NULL, group=NULL, weight=NULL), person.grouping=NULL, item.grouping=NULL, model.statement="item", m.model="1pl", Title = NULL, jobName, jobFolder, subFolder=list(), dataName=NULL, anchor=NULL, pathConquest, method=NULL, std.err=NULL, distribution=NULL, n.plausible=NULL, set.constraints=NULL, nodes=NULL, p.nodes=NULL, f.nodes=NULL, n.iterations=NULL, converge=NULL, deviancechange=NULL, name.unidim=NULL, equivalence.table="wle",use.letters=FALSE, check.for.linking=FALSE)
```

Arguments

dataset data.frame containing all variables necessary for analysis

ID name or column number of 'id' variable

regression character vector with names or integer vector with column numbers of one or

more 'context' variables (e.g. sex, school, ...)

DIF character string with name or scalar with column number of only one variable

denominating groups for which analysis of differential item functioning is to be

done (e.g. sex, class, ...)

group.var character vector with names or integer vector with column numbers of one or

more 'group' variables (e.g. sex, school, ...)

weight character string with name or scalar with column number of only one 'weight-

ing' variable

testitems character vector with names or integer vector with column numbers of 'item'

variables (e.g. sex, school, ...)

na List of numerical vector including numbers to be considered as 'sysmis'. Spe-

cific missing codes can be defined for each type of variable, e.g. testitems, DIF

variables, ...

item.grouping data.frame with grouping information of items, first column must be 'item'

which includes item names, further columns contain scale definitions, 0 indicates that the respective item is NOT part of the scale, 1 indicates that this item

is part of the scale, colnames of columns are the names of the scales

person.grouping

data.frame with grouping information of persons, first column must be the name of 'id' (e.g. idstud), further columns contain group definitions, 0 indicates that the respective person is NOT part of the group, 1 indicates that this person is

part of the group, colnames of columns are the names of the groups

model.statement

character string which appears in Conquest Syntax as model statement. Set to item by default. When DIF variable is specified, statement is set to item - [name of DIF variable] + item*[name of DIF variable] by default. However, user's specification of model.statement overwrites default in each

case.

m.model character string specifying the IRT model used for analysis. At the time, only

"1PL" is available.

Title optional: character string with title of analysis which appears in Conquest Syn-

tax. If no title is specified, informations about computer and user name and R

version are printed in Conquest Syntax.

jobName character string specifying name of analysis. All Conquest input and output files

will named by jobName and their corresponding suffixes.

jobFolder character string specifying the folder where all analysis files will be generated,

for example "C:/programme/analysis"

subFolder optional: List of character strings specifying maximal 2 optional subfolders rel-

ative to jobFolder for the data and the output files. Character strings must be

named with data and out, for example subFolder=list(data="../../dataset/analysis1",

out="../../output/analysis1") Double dots . . indicates to move one level above in folder structure. For example, if jobFolder is "C:/programme/analysis"

and subFolder is list(data="../dataset/analysis1", out="../../output/analysis1"),

dataset is written to "C:/programme/dataset/analysis1" and output is written to "C:/output/analysis1". When subFolder\$data == NULL, dataset is written to the folder specified by jobFolder. Same is true for subFolder\$out == NULL.

dataName optional: character string specifying name of dataset if intend to differ from

name specified by jobName. When dataName == NULL, dataset is named [jobName]. dat

anchor optional: data frame with anchor parameters. First column of anchor refers to

item name, second columns refers to parameter used as reference for anchoring.

pathConquest character string with path and name of Conquest console, for example "c:/programme/conquest/co

method optional: character string with method for analysis. Possible options are "gauss"

(default), "quadrature", "montecarlo". See Conquest manual, pp.225.

std.err optional: character string specifying types of standard errors to be estimated.

Possible options are "full", "quick" (default), "none". See Conquest manual,

pp.167.

optional: character string with a priori trait distribution. Possible options are distribution

"normal" (default), "discrete". See Conquest manual, pp.167.

optional: integer scalar specifying numbers of plausible values to draw. Default n.plausible

set.constraints

optional: character string specifying the constraints of the scale. Possible options are "cases" (default), "items", "none". When anchor parameter are

specified in anchor, constraints will set to "none" in each case.

nodes optional: integer scalar specifying numbers of nodes in analysis. Default is 15.

optional: integer scalar specifying numbers of p nodes in analysis. Sets the p.nodes

number of nodes that are used in the approximation of the posterior distributions, which are used in the drawing of plausible values and in the calculation of EAP

estimates. The default is 2000.

f.nodes optional: integer scalar specifying numbers of f nodes in analysis. Sets the

number of nodes that are used in the approximation of the posterior distributions

in the calculation of fit statistics. The default is 2000.

optional integer scalar. Sets the maximum number of iterations for which estin.iterations

mation will proceed without improvement in the deviance. The minimum value

permitted is 5. The default value is 20.

optional scalar. Instructs estimation to terminate when the largest change in any converge

parameter estimate between successive iterations of the EM algorithm is less

than converge. The default value is 0.0001.

optional scalar. Instructs estimation to terminate when the change in the dedeviancechange

viance between successive iterations of the EM algorithm is less than deviancechange.

The default value is 0.0001.

name.unidim optional: character string with name of one latent dimension, if not specified in

labels.

equivalence.table

optional: character string specifying type of equivalence table to print. Possible

options are "wle" (default), "mle" or NULL.

use.letters logical: Should values coded als letters? May be relevant only in partial credit

models comprising items with more than 10 categories to avoid columns with

width 2 in Conquest.

check.for.linking

logical: If TRUE, items in dataset are checked for being connected with each other via design.

Value

No results are returned to console. Input files and batch string are written to disk in specified folder(s).

Author(s)

Sebastian Weirich, Karoline Sachse, Martin Hecht

automateDataPreparation

automateDataPreparation

Description

prepare datasets for automateModels

Usage

```
automateDataPreparation( inputDat = NULL, inputList, path = NULL, loadSav,
checkData, mergeData, recodeData,aggregateData, scoreData, writeSpss,
filedat = "zkddata.txt", filesps = "readZkdData.sps",
aggregatemissings = "use.default", rename = TRUE, recodedData = TRUE,
correctDigits=FALSE, truncateSpaceChar = TRUE, newID = NULL, oldIDs = NULL,
missing.rule = list(mvi=0, mnr=0, mci=0, mbd=NA, mir=0, mbi=0))
```

Arguments

| inputDat | A list of data frames if no .sav files shall be read in. |
|---------------|--|
| inputList | A list of data frames containing additional information (see Details). |
| path | A character string containing the path where the logfolder will be created. Also required by loadSav (source of SPSS files) and writeSpss. Default is the current R working directory. |
| loadSav | logical (whether function loadSav shall be called). |
| checkData | logical (whether function checkData shall be called). |
| mergeData | logical (whether function mergeData shall be called). |
| recodeData | logical (whether function recodeData shall be called for subunits). |
| aggregateData | logical (whether function aggregateData shall be called). |
| scoreData | logical (whether function recodeData shall be called for units). |
| writeSpss | logical (whether function writeSpss shall be called). |
| 0.7 | |

filedat A character string with the name of the output data file required by writeSpss.

filesps A character string with the name of the output syntax file required by writeSpss.

missing.rule A list containing recode information for character missings required by writeSpss.

See 'References' for description of default values.

 ${\tt aggregatemissings}$

A character string. Either "use.default" or "seeInputList", if pattern was

specified in inputList\$aggrMiss.

rename logical. See aggregateData.

automateModels 9

Details

inputList is a list of data frames. It can be created either by ZKDaemon via readDaemonXlsx or by makeInputLists. Compulsory: units, subunits, values. Optional: unitRecodings, savFiles, newID, aggregateMissings.

Value

A single data frame in last transformation status.

Author(s)

Karoline Sachse

References

http://code.google.com/p/zkdlib/wiki/MissingHandling

Examples

automateModels

automateModels

Description

specify and run several ConQuest models

Usage

```
automateModels(dataset, id = NULL, context.vars = NULL, items = NULL,
item.grouping = NULL, select.item.group = NULL, person.grouping.vars = NULL,
person.grouping.vars.include.all = FALSE, person.grouping = NULL,
select.person.group = NULL, additional.item.props = NULL, folder,
overwrite.folder = TRUE, analyse.name.prefix = NULL, analyse.name = NULL,
analyse.name.elements = NULL, data.name = NULL, m.model = NULL, software = NULL,
dif = NULL, weight = NULL, anchor = NULL, regression = NULL,
```

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```
adjust.for.regression = FALSE, q3 = FALSE, missing.rule = NULL, cross = NULL,
subfolder.order = NULL, subfolder.mode = NULL, allNAdelete = TRUE, additionalSubFolder = NULL,
run.mode = NULL, n.batches = NULL, run.timeout = 1440, run.status.refresh = 0.2,
all.local.cores = TRUE, email = NULL, smtpServer = NULL, write.txt.dataset = FALSE,
delete.folder.countdown = 5, conquestParameters = NULL )
```

Arguments

dataset data.frame containing all variables type of variables ("id", "context.vars" or

"items") must be set using options id, context.vars, items

id name or column number of 'id' variable in dataset

context.vars names or column numbers of 'context' variables (e.g. sex, school, ...) in

dataset

items names or column numbers of 'item' variables in dataset if omitted, all variables

that are not classified as 'id' or 'context' variables are treated as 'items'

item.grouping data.frame with grouping information of items, first column must be 'item'

which includes item names, further columns contain scale definitions, 0 indicates that the respective item is NOT part of the scale, 1 indicates that this item

is part of the scale, colnames of columns are the names of the scales

select.item.group

character vector of scale names chosen for analysis

person.grouping.vars

character vector of 'context' variables in dataset which are used to automatically generate 'person.grouping', each category is transformed into the 'per-

son.grouping' format

person.grouping.vars.include.all

logical vector (along person.grouping.vars), indicates whether to generate a variable 'all' for the specific variable

person.grouping

data.frame with grouping information of persons, first column must be the name of 'id' (e.g. idstud), further columns contain group definitions, 0 indicates that the respective person is NOT part of the group, 1 indicates that this person is part of the group, columns of columns are the names of the groups

select.person.group

character vector of group names chosen for analysis

additional.item.props

data.frame of additional item information to be merged to model results, first column must be 'item' and contain item names

folder to write output into

folder folder overwrite.folder

logical, if TRUE (default), folder is completely emptied

analyse.name.prefix

prefix (e.g. "pilotStudy") to be attached to all analyses names

analyse.name analyses names are usually automatically set, if you want to set them manually

use this option

analyse.name.elements

analyses names are set automatically using these elements: c ("scale" , "group" , "dif" , "regression" , "anchor"), use this option to change composition and order of the analyses names generation

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data.name optional: character string specifying name of dataset if intend to differ from

name specified by jobName. When dataName == NULL, dataset is named [job-

Name].dat

m.model measurement model, "1pl" (default), "2pl", "3pl", "4pl"

software "conquest" (default) no other software implemented yet

dif variable that is used for differential item functioning

weight case weight variable

anchor data.frame with anchor information

regression variable(s) that is/are used

adjust.for.regression

if TRUE item parameters (difficulty) are centered on the mean of the entire sample if FALSE (default) item parameters (difficulty) are centered on the mean of

the regression reference group

q3 Logical: If TRUE, Yen's Q3 statistic is computed.

missing.rule definition how to recode distinct missings in dataset

cross scales in 'item.grouping' and groups in 'person.grouping' can be crossed to de-

fine distinct analyses "all": scales and groups are crossed "item.groups", scales are separately (unidimensional) run (instead of one multidimensional model) "person.groups", person groups are separately (single group) run (instead of one

multigroup model)

subfolder.order

subfolders are automatically generated in this order c ($"i.model"\ ,\ "p.model"\ ,$

"m.model", "software", "dif", "regression", "anchor")

subfolder.mode "none": no subfolders are created "full": complete subfolders are created ac-

cording to 'subfolder.order' "intelligent" (default): meaningful subfolders are

created

allNAdelete if TRUE all cases with complete missings on items are removed, if FALSE these

cases are not deleted Note: this is a global option, that is set for all modelss

additionalSubFolder

specification for 'data' and 'out' subfolder (constant over all analyses)

run.mode "serial": serial runs on local machine "parallel": batch files must be started man-

ually (e.g. on several machines)

n. batches number of batch files that are created, batch files contain one or more analyses

run.timeout minutes to wait for analyses to finish, default: 1440 (24h)

run.status.refresh

time for console refresh of model run status, default: 0.2 (12sec)

all.local.cores

if TRUE and run.mode="serial" all cores of local machine are used for analysis

email set email address to receive an email when analyses are finished or time's up

smtpServer for sending emails, default: "mailhost.cms.hu-berlin.de"

write.txt.dataset

write out datasets as ascii, default: FALSE

delete.folder.countdown

countdown for deletion of 'folder', default: 5 (seconds)

12 bi.linking

conquestParameters

Set ConQuest parameters as a named list.

Available option are:

"pathConquest", "method", "std.err", "distribution", "n.plausible", "set.constraints", "nodes", "p.nodes", "f.nodes", "n.iterations", "converge", "deviancechange", "equivalence.table", "use.letters", "na", "model.statement"

See automateConquestModel documentation for details.

Details

To run several models list parameters as corresponding lists Explicitly list NULL if parameter should not be set or be defaulted See examples

Value

returns results in specific format

Author(s)

Martin Hecht, Karoline Sachse, Sebastian Weirich, Christiane Penk, Malte Jansen, Sebastian Wurster

bi.linking bi.linking

Description

Links results from several analysis. Each analysis is linked with each other.

Usage

```
bi.linking \ (\ results \ ,\ scales=NULL \ ,\ folder=NULL \ ,\ file.name=NULL \ ,\ method = NULL \ ,\ lower.trianger \ )
```

Arguments

results result list from automateModels run

scales Character vector of scales for which linking should separately done. If NULL, all analysis in the results list are linked. Note: due to suboptimalities in development process, analysis name must contain scale!!

folder output folder, will be emptied!

file.name for output excel, default: "bi.linking.results.xlsx"

method set linking method to either "Mean-Mean", "Haebara" or "Stocking-Lord" (de-

fault)

lower.triangle set reference groups for the linking

Value

writes linking results to excel file. returns linking results as list.

Author(s)

Martin Hecht

checkData 13

| checkData | Check Datasets for Missing Values and Invalid Codes | |
|-----------|---|--|
| | | |

Description

Check data frames for missing or duplicated entries in the ID variable, persons and/or variables without valid codes, and invalid codes. Invalid codes are codes which are not specified in table values.

Usage

```
checkData (dat, values, subunits, units)
```

Arguments

| dat | A data frame |
|----------|---|
| values | A data frame with code information. See 'Details'. |
| subunits | A data frame with subunit information. See 'Details'. |
| units | A data frame with unit information. See 'Details'. |

Details

The results of checkData will be written to a protocol file with sunk.

Examples of data frames values, subunits and units can be found via data(inputList).

Value

Used for its side effects. The return value is NULL.

Author(s)

Nicole Haag, Anna Lenski

References

For missing types see http://code.google.com/p/zkdlib/wiki/MissingHandling

See Also

sunk

14 checkInput

| checkInput | Check Input Data Frames | |
|------------|-------------------------|--|
| | | |

Description

Check input data frames for consistency and replace missing information with default values (if necessary).

Usage

```
checkInput(values, subunits, units, checkValues = TRUE, checkUnits = TRUE)
```

Arguments

| values | A data frame with code information. See 'Details' |
|-------------|--|
| subunits | A data frame with subunit information. See 'Details' |
| units | A data frame with unit information. See 'Details'. |
| checkValues | Logical: Should data frame values be checked? |
| checkUnits | Logical: Should data frame units be checked? |

Details

This function is largely for internal use and is called by makeInputLists before lists are generated. Examples of data frames values, subunits and units can be found via data(inputList).

Value

A list containing the checked and (if necessary) defaulted input data frames:

| values | Checked data frame with code information. Will be returned if checkValues = TRUE. |
|----------|---|
| subunits | A data frame with subunit information. |

A data frame with unit information. Will be returned if checkUnits = TRUE.

Warning

units

Function will not check input data frames if checkValues and checkUnits are both FALSE.

Author(s)

Nicole Haag

See Also

makeInputLists

checkLink 15

Description

checks whether items in a dataset are linked via design

Usage

```
checkLink ( dataFrame, sysmis = NA, verbose = TRUE)
```

Arguments

dataFrame A data.frame where all columns denote test items

sysmis character string specifying values to be treat as missing by design

verbose logical: Should output printed to console?

Value

A logical value, i.e. TRUE or FALSE, indicating whether items in dataset are linked to each other.

Author(s)

Sebastian Weirich

collapseMissings Collapse Missings

Description

converts character missings of different types to 0 or NA

Usage

```
collapseMissings(dat, missing.rule = NULL, item.names)
```

Arguments

data frame containing character missings (e.g. type 'mbd' - missing by design)
missing.rule list, definition how to recode distinct missings in dataset. See details for default.
item.names character vector containing column names of the data frames whose character missings are to be collapsed.

missings are to be collapsed

Details

Default missing.rule in collapseMissings is: text volume insufficient = 0, missing not reached = 0, missing coding impossible = NA, missing by design = NA, missing invalid response = 0, missing by intention = 0

The results of collapseMissings will be written to a protocol file with sunk.

16 commonItems

Value

A data frame with recoded missings.

Author(s)

Karoline Sachse, Martin Hecht

References

For missing types see http://code.google.com/p/zkdlib/wiki/MissingHandling

Examples

```
data(inputDat)
dat1 <- inputDat[[1]] # get first dataset from inputDat
datColMis <- collapseMissings(dat = dat1,
missing.rule = list(mvi = 0 ,mnr = 0 ,mci = 0 ,mbd = NA ,mir = 0 ,mbi = 0),
item.names=colnames(dat1)[- c(1:2)])</pre>
```

commonItems

identify common items of groups

Description

This function identifies items that groups of persons have in common.

Usage

```
commonItems ( data , group.var , missing = NA , uncommon = FALSE , simplify = TRUE )
```

Arguments

data data.frame group.var group variab

group variable in data.frame, eihter numeric indicator of column or column

name

missing missing specification

uncommon if TRUE a vector of uncommon items is additionally returned

simplify if TRUE a character vector is returned (only in case of 2 groups and uncom-

mon=FALSE)

Value

returns a list of all group.var combinations with character vectors of common item names if uncommon=TRUE a vector of uncommon (unique) items of each group is additionally returned names of list are both group names concatenated by "I"

Author(s)

Martin Hecht

crop 17

Examples

```
data(science1)
d <- science1[,c("version",science1.items)]

# common items ar listed for each combination of groups
str ( commonItems ( data = d , group.var = "version" , missing = "mbd" ) )

# uncommon items are returned as well
str ( commonItems ( data = d , group.var = "version" , missing = "mbd" , uncommon = TRUE ) )

crop crop</pre>
```

Description

remove trailing and leading characters from character strings

Usage

```
crop ( x , char = " ")
```

Arguments

x character string

char character to be removed from beginning and end of x

Author(s)

Martin Hecht, Sebastian Weirich

detect.suppression detect suppression effects in regression models

Description

This function detects suppression effects in regression models.

Usage

```
detect.suppression ( data , dependent , independent , full.return = FALSE , xlsx.path = NULL )
```

Arguments

| data | data.frame with data to be used |
|-------------|--|
| dependent | dependent variable in regression model |
| independent | character vector of independent variables in regression model |
| full.return | if FALSE a data.frame as a quadratic matrix with suppression effects (TRUE/FALSE) of independent variables is returned |
| | if TRUE a data.frame with all calculated terms ist returned |
| xlsx.path | full path of Excel file that results should be written to |

18 detect.suppression

Details

formulae (13.39a) and (13.39b) decribed in Bortz (1999) page 446 are used

if full.return=TRUE a data.frame is returned.

Columns are:

rownames: <dependent variable> ~ <independent variables> | <independent variable that is tested for suppression>

multiple.reg: logical, indicates wether there are 2 (FALSE) or more than 2 (TRUE) independent variables in the regression model

dep: dependent variabel in regression model

pred: independent variable that is investigated on suppression effect

preds: independent variables in regression model besides pred

cor_pred_c: correlation of pred and dependent variable

cor_pred_fitted_c: correlation of predicted pred by indepenent variables and dependent variable

r.sq_pred: R squared from model predicting pred by independent variables

rterm.minus: right term in formula (13.39a) rterm.plus: right term in formula (13.39b)

rterm.minus.diff: difference of rterm.minus and cor_pred_c

rterm.plus.diff: difference of cor_pred_c and rterm.plus

(positive difference of rterm.minus.diff or rterm.plus.diff indicates suppression effect)

rterm.minus.log: logical value of formula (13.39a) rterm.plus.log: logical value of formula (13.39b)

suppression: logical, rterm.minus.log|rterm.plus.log

if full.return=FALSE a data.frame as quadratic matrix is returned:

rows and columns are independent variables

 $diagonal\ includes\ suppression\ for\ suppression\ effect\ of\ variable\ in\ multiple\ regression$

triangles include suppression for bivariate independent variables, "row" suppresses "column"

Value

depends on options full.return

Author(s)

Martin Hecht

References

for formulae used by detect.suppression see

Bortz, J. (1999). Statistik fuer Sozialwissenschaftler. 5. Auflage. Berlin: Springer. p. 446

dichotomize 19

dichotomize

dichotomize a numeric vector

Description

dichotomize a numeric vector by median or mean split

Usage

```
dichotomize ( numvec , method = c("median", "mean") , randomize = TRUE , ... )
```

Arguments

numvec numeric vector

method either median or mean split

randomize logical, if TRUE elements that equal the split threshold are randomly assigned to

one of the two groups if FALSE default behavior of cut is used

... arguments are passed to set.seed and cut

Value

returns vector with dichotomization indicators

Author(s)

Martin Hecht

Examples

```
numvec <- c(1,2,3,4,5)
dichotomize ( numvec )

# set seed for random assignment of elements that match split threshold by passing argument 'seed' to function
# ( '3' in numvec is on threshold if median is used )
dichotomize ( numvec , seed = 12345 )

# set level names by passing argument 'labels' to cut function
dichotomize ( numvec , labels = c ( "low" , "high") )</pre>
```

 ${\tt exploreDesign}$

explore data design

Description

explore data structure with respect to specific missing code (e.g. "missing by design")

Usage

```
{\tt exploreDesign~(~data~,~missing~=~NA~,~id~=~NULL~,~itemsPerPerson~=~TRUE~,~personsPerItem~=~TRUE}
```

20 get.dsc

Arguments

data data.frame

missing missing specification
id variable in data if exists

itemsPerPerson logical, if TRUE items per person list is returned personsPerItem logical, if TRUE persons per item list is returned

Value

depends on itemsPerPerson and personsPerItem , if both are TRUE a list with both elements is returned

Author(s)

Martin Hecht

Examples

```
data(science1)
d <- science1[,!colnames(science1) %in% science1.context.vars]
design <- exploreDesign ( data = d , missing = "mbd" , id = "id" )
str(design)</pre>
```

get.dsc

Reads Conquest descriptive files.

Description

Reads Conquest files comprising descriptive population statistics generated by the 'descriptives' statement.

Usage

```
get.dsc(file)
```

Arguments

file

Character string of the Conquest descriptives file.

Value

A list of n elements, with n the number of groups in the analysis. Each element is a list with two data frames, the single and aggregated descriptives of the corresponding group. Single descriptives give for each dimension the number of observations, mean, standard deviation and variance of the corresponding estimate, i.e. the WLE or the plausible values (PVs). When descriptives for PVs are read in, mean, standard deviation and variance for each pausible value is given. Aggregated descriptives give mean, standard deviation and variance of the standard error of corresponding estimate. When descriptives vor plausible values are read in, aggregated descriptives gibt also mean, standard deviation and variance of all plausible values.

get.equ 21

References

See Conquest manual, pp.162.

get.equ

Reads equivalence table created in Conquest analysis.

Description

Reads Conquest files comprising equivalence tables for MLE or WLE parameters.

Usage

```
get.equ(file.equ)
```

Arguments

file.equ

Character string of the Conquest equ-file.

Value

A list of n+1 elements, with n the number of dimensions in the analysis. Each element is a data.frame, whose name correponds to the name of the dimension the values belongs to. All data.frames except the last one give the transformation of each possible raw score to the WLE or MLE score including it's standard error. First column in each data.frame contains the raw score, second column the transformed WLE or MLE score, third columns it's standard error.

The last element of the list give some sparse information about the model specifications.

References

See Conquest manual, pp.162.

get.itn

get.itn

Description

blablabla

Usage

get.itn(file)

Arguments

file

22 get.q3

| get.plausible | Reads Conquest plausible values files |
|---------------|---------------------------------------|
| | |

Description

Function reads Conquest plausible value files and transforms them into a R data frame.

Usage

```
get.plausible(file)
```

Arguments

file

Character string of the Conquest plausible values file to be read in.

Details

Funktion identifies number of cases, number of plausible values and number of dimensions.

Value

A data frame, where each row corresponds to one case. Columns are labeled with dimension names and number of corresponding plausible value.

| case | Case number. Each row represents one person. |
|--------|---|
| ID | Case ID, if listed in Conquest plausible values file. |
| pv | Plausible value. Denotation of columns names is pv.[name of dimension]_[number of plausible value]. For example, pv.reading_6 refers to the 6th plausible value of reading dimension. |
| eap | Expectation value of the a posterio distribution of the corresponding dimension. |
| eap.se | Standard error of the EAP estimate. |
| get.q3 | get.q3 |

Description

```
get Q3 statistics
```

Usage

```
get.q3 ( results )
```

Arguments

results

results (structured list) from automateModels run

Value

list (analyses) of data.frames in matrix format containing Q3 statistics

get.shw 23

Author(s)

Martin Hecht

| get.shw | Reads Conquest showfiles | |
|---------|--------------------------|--|
| | | |

Description

Function reads Conquest showfiles and transforms them into a R list of data frames.

Usage

Arguments

| file | Character string of the Conquest showfile to be read in. |
|---------------|--|
| dif.term | Optional: Character string. Name of the term considered to be DIF-term. Must match corresponding term in showfile. |
| split.dif | Logical: When TRUE, DIF-Parameter are only given for Reference group. |
| abs.dif.bound | When DIF-Parameter are evaluated, this specifies the critical value for absolute DIF. |
| sig.dif.bound | When DIF-Parameter are evaluated, this specifies the critical value for confidence interval DIF. |

Details

Funktion searches for 'TERM'-statements in Conquest showfile and reads the tables associated with. If one statement is specified to contain DIF analyses, absolute DIF value is computed 2*[group-specific parameter]. Confidence intervalls for 90, 95 and 99 percent are computed via the standard error of specific parameters. If both criteria - absolute DIF exceeds abs.dif.bound and the confidence intervall does not include sig.dif.bound, item is considered to have DIF.

Value

A list of data frames, named by the 'TERM'-statements in Conquest showfile, plus an additional data frame with regression coefficients when latent linear regression model was specified in Conquest analysis. If one term was specified as DIF-statement, the corresponding data frame is augmented with additional columns for confidence intervals and indicators specifying significant DIF.

Each data frame corresponding to a 'TERM' statement contains following columns:

| item | Name of item |
|----------|---|
| ESTIMATE | Estimated difficulty of item |
| ERROR | Standard error of estimated item difficulty |
| MNSQ | Item's 'Outfit' |
| MNSQ.1 | Items's 'Infit' |
| CI | Lower and upper bound confidence intervals |

24 get.wle

| T | T values, corresponding to confidence intervals |
|----------|--|
| filename | Name of show file read in |
| abs.dif | Only for DIF analysis. Absolute DIF, computed as $2*[group-specific parameter]$. |
| ci.lb | Lower bound confidence interval for specific significance level of 90, 95 or 99 percent. |
| ci.ub | Upper bound confidence interval for specific significance level of 90, 95 or 99 percent. |
| sig | Indicates whether the corresponding item matches both DIF criteria. See details. |

When latent regression was specified, the last element of the returned list is a data frame with regression coefficients, corresponding to the number of dimensions and the number of regressors. Regressor names, regression coefficients and its standard errors are given for each dimension.

Rows represent the regressors, columns represent the latent dimension to which the regression is fitted.

| get.wle | Reads Conquest WLE or MLE files. | |
|---------|----------------------------------|--|
| | | |

Description

Reads Conquest files comprising maximum likelihood estimates (MLE) or weighted likelihood estimates (WLE).

Usage

```
get.wle(file)
```

Arguments

file Character string of the Conquest MLE or WLE file to be read in.

Value

A data frame with columns according to the corresponding MLE or WLE file. For each dimension of the analysis number of solved items, number of presented items, point estimate and its standard error is given. Each row represents one person. Columns are named as follows:

| case | Case number. Each row represents one person. |
|----------|---|
| n.solved | Number of solved items by the i-th person. |
| n.total | Number of total items presented to the i-th person. |
| wle | WLE or MLE estimate. |
| wle.se | Standard error of WLE or MLE estimate. |

The last number of columns names represents the dimension the WLE or MLE estimate belongs to.

inputDat 25

inputDat

List of Three Datasets from Educational Assessment

Description

Simulated data for three booklets for an educational assessment study.

Usage

```
data(inputDat)
```

Format

This list contains 3 data frames, each with the following columns:

ID Person-ID

Hisei A continuous covariate.

Ixx Item responses to a selection of 30 test items.

Details

code, subunit and unit descriptions are stored in dataset inputList.

Examples

```
data(inputDat)
str(inputDat)
```

inputList

Data Frames with Code, Subunit and Unit Information for Datasets in inputDat

Description

These data frames contain information about codes, subunits and units for the datasets in inputDat and are necessary inputs for functions automateDataPreparation, checkData, recodeData and aggregateData.

Usage

```
data(inputList)
```

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Format

A list with three data frames:

1. units: Unit information, contains the following columns:

unit Unit name.

unitType Subunit types: ID = ID variable; TI = test item; CV = context variable.

unitLabel Unit label, to be used by writeSpss.

unitDescription Unit description.

unitAggregateRule Aggregate rule for unit: SUM; MEAN.

unitScoreRule Scoring rule for unit (not sure how this will be used in the future.)

2. subunits: Subunit information, contains the following columns:

unit Unit name, for which subunits are given.

subunit Subunit name.

subunitType Subunit types:'?'.

subunitLabel Subunit label, to be used by writeSpss.

subunitDescription Subunit descriptions.

subunitPosition Subunit position in test booklet (e.g., line 1).

subunitTransniveau Subunit transformation level.

subunitRecoded Name of recoded subunit.

subunitLabelRecoded Label for recoded subunit, to be used when writeSpss is applied to a dataset produced by recodeData.

3. values: Value information, contains the following columns:

subunit Subunit name, for which values are given.

value Valid values for the respective subunit.

valueRecode Recode values for the respective value.

valueType Value types: vc = valid code; mbd = missing - by design; mvi = missing - volume insufficient; mnr = missing - not reached; mci = missing - coding impossible; mbi = missing - by intention.

valueLabel Value labels, to be used by writeSpss.

valueDescription Value descriptions.

valueLabelRecoded Labels for recoded values, to be used when writeSpss is applied to a dataset produced by recodeData.

valueDescriptionRecoded Descriptions for recoded values.

4. unitRecodings: Unit recoding information, contains the following columns:

unit Unit name

value Valid values for the respective unit.

valueRecode Recode values for the respective value.

valueType Value types: vc = valid code; mbd = missing - by design; mvi = missing - volume insufficient; mnr = missing - not reached; mci = missing - coding impossible; mbi = missing - by intention.

valueLabel Value labels, to be used by writeSpss.

valueDescription Value descriptions.

valueLabelRecoded Labels for recoded values, to be used when writeSpss is applied to a
 dataset produced by recodeData.

5. savFiles: information for loadSav, contains the following columns:

loadSav 27

```
filename SPSS filenames
```

case.id ID variable in the respective dataset, used by mergeData

6. newID: information for mergeData, contains the following columns:

key one of the entries should be master-id

value the corresponding value; how the ID variable in the final dataset shall be named

7. aggrMiss: missing aggregation pattern for aggregateData

Examples

```
data(inputList)
str(inputList)
```

loadSav

loadSav

Description

read SPSS data files and change id names, if necessary

Usage

```
loadSav(path = getwd(), savFiles = NULL, oldIDS, newID, correctDigits = FALSE, truncateSpaceChar
```

Arguments

```
path
savFiles
oldIDS
newID
correctDigits
truncateSpaceChar
```

Examples

```
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##--or do help(data=index) for the standard data sets.

## The function is currently defined as
function ( path=getwd(), savFiles=NULL, oldIDS, newID, correctDigits=FALSE, truncateSpaceChar = TRUE ) {
    funVersion <- "loadSAV_0.0.2"
    if(missing(oldIDS)) {stop(paste("Error in ",funVersion,": 'oldIDS' is missing.\n",sep="")) }
    if(missing(newID)) {stop(paste("Error in ",funVersion,": 'newID' is missing.\n",sep="")) }
    if(length(newID)!=1) {stop(paste("Error in ",funVersion,": 'newID' has to be of length 1.\n",sep # if(!exists("read.spss")) {library(foreign)}
    if(!is.null(savFiles)) {
        fileExists <- file.exists(file.path(path,savFiles))
        if(all(!fileExists)) {
            stop(paste("Error in ",funVersion,": None of the files specified in 'savFiles' were found</pre>
```

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```
if(!all(fileExists)) {
               cat(paste(funVersion,": Following files specified in 'savFiles' were not found in ",path,
               notFoundFiles <- savFiles[!fileExists]</pre>
               FoundFiles
                                          <- savFiles[fileExists]</pre>
               cat(paste(notFoundFiles,collapse=", "))
               cat("\n0nly found files will be read in.\n")
               savFiles
                                             <- savFiles[fileExists]</pre>
        }
 if(is.null(savFiles)) {
        savFiles <- list.files(path=path,pattern=".sav|.SAV",recursive=FALSE)</pre>
        if(length(savFiles)==0) {
               stop(paste("No '.sav'-files found in ",path,".\n",sep=""))
 cat(paste(funVersion,": Found ", length(savFiles), " 'savFiles' in ",path,".\n",sep=""))
 ### hier beginnt das eigentliche Einlesen
 allDataFrames <- NULL
  for (i in seq(along=savFiles)) {
             file.i <- data.frame(read.spss(file.path(path,savFiles[i]),to.data.frame=FALSE, use.value.</pre>
             idCol <- unique(unlist(lapply(oldIDS, FUN=function(ii) {grep(ii,colnames(file.i))})))</pre>
             if(length(idCol)<1) {</pre>
                   stop(paste("Error in ",funVersion,": None of the specified 'oldIDS' were found in datase
             if(length(idCol)>1) {
                   stop(paste("Error in ",funVersion,": More than one of the specified 'oldIDS' were found
            colnames(file.i)[idCol] <- newID</pre>
             ### Leerzeichen abschnipseln
             if(truncateSpaceChar == TRUE) {
                   for (ii in 1:ncol(file.i)) {
                              file.i[,ii] <- crop(file.i[,ii])</pre>
             }
             ### Stelligkeitskorrektur
             if(correctDigits == TRUE) {
                   colsToCorrect <- lapply(1:ncol(file.i), FUN=function(ii) { sort(unique(nchar(file.i[,ii])</pre>
                   options(warn = -1)
                   colsToCorrect <- which( unlist( lapply(colsToCorrect, FUN=function(ii) { all(ii == c(1))</pre>
                   options(warn = 0)
                   if(length(colsToCorrect)>0) {
                          \verb|cat(paste(funVersion,": ",length(colsToCorrect)," columns are corrected for column with the column of the colu
                         for (ii in colsToCorrect) {
                                     file.i[,ii] <- gsub(" ","0", formatC(as.character(file.i[,ii]),width=2))</pre>
                   }
             allDataFrames[[i]] <- file.i
 }
return(allDataFrames)
```

long2matrix 29

Description

transforms long format data.frame into a matrix format data.frame

Usage

```
long2matrix ( data , sort = TRUE , triangle = NULL ,
force.diagonal = FALSE , exclude.diagonal = FALSE ,
long2matrix = TRUE )
```

Arguments

```
data.frame with columns "row", "col", "val"

sort sort rows and columns of matrix

triangle if not NULL a symmetric matrix will be constructed available options are "upper", "lower", "both"

force.diagonal a diagonal is forced into matrix even if no diagonal elements are in data exclude.diagonal

the diagonal is excluded if possible

long2matrix if FALSE data is not transformed
```

Value

Author(s)

Martin Hecht

Examples

```
d1 <- data.frame (
"row" = c ( "v1" , "v2" , "v2" , "v3" , "v1" , "v3" ) ,
"col" = c ( "v1" , "v3" , "v2" , "v1" , "v2" , "v3" ) ,
"val" = c ( 1 , 5 , 4 , 3 , 2 , 6 ) , stringsAsFactors = FALSE )
# unsorted matrix
long2matrix ( data = d1 , sort = FALSE )
# sorted by default
long2matrix ( data = d1 )
# extract upper triangle of symmetric matrix
long2matrix ( data = d1 , triangle = "upper" )
# exclude diagonal elements
long2matrix ( data = d1 , triangle = "upper" , exclude.diagonal = TRUE )
# if full matrix ("both" triangles) is requested, the diagonal cannot be excluded, option is ignored
long2matrix ( data = d1 , triangle = "both" , exclude.diagonal = TRUE )
# no diagonal elements are specified
d2 <- data.frame (
"row" = c ( "v2" , "v1" , "v1" ) ,
"col" = c ( "v3" , "v3" , "v2" ) ,
"val" = c ( 5 , 3 , 2 ) , stringsAsFactors = FALSE )
```

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```
long2matrix ( data = d2 )
# diagonal is set (with NAs)
long2matrix ( data = d2 , triangle = "upper" , force.diagonal = TRUE )
```

makeCodebookInput

Make Input Data Frames From IQB-Codebooks

Description

Make Input Data Frames From IQB-Codebooks

Usage

```
makeCodebookInput(codebook)
```

Arguments

codebook

dataframe IQB-Codebook

Details

xxx

Value

XXX

makeInputLists

 $\begin{tabular}{ll} \textit{Generate Input Lists for Functions} & \texttt{checkData}, & \texttt{recodeData} & \texttt{and} \\ \texttt{aggregateData} & \end{tabular}$

Description

Transforms information given in values, subunits and units in a format that is used by checkData, recodeData and aggregateData.

Usage

```
makeInputLists(values, subunits, units, recodedData = TRUE)
makeInputCheckData(values, subunits, units)
makeInputRecodeData(values, subunits)
makeInputAggregateData(subunits, units, recodedData = TRUE)
```

Arguments

values A data frame with code information. See Details.
subunits A data frame with subunit information. See Details.
units A data frame with unit information. See Details.

names (as in subunits\$subunit) or recoded subunit names (as in subunits\$subunitRecoded).

Default is TRUE, meaning that colnames are recoded subitem names. This pa-

rameter is only relevant when input for aggregateData is generated.

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Details

This function generates specific inputs for the data preparation functions checkData, recodeData and aggregateData. It is largely for internal use of these functions, who call their respective version.

Examples of data frames values, subunits and units can be found via data(inputLists).

Value

A list with several of the following entries (depending on which version of the function is called):

varinfoRaw A list with information about variables and their values expected in raw data.

varinfoRecoded A list with information about variables and their values expected in recoded

data

varinfoAggregated

A list with information about variables and their values expected in aggregated

data.

recodeinfo A list with information needed for recoding of data.

aggregateinfo A list with information needed for aggregation of data.

Author(s)

Nicole Haag

Examples

```
data(inputList)
lists <- makeInputLists(inputList$values, inputList$subunits, inputList$units, recodedData = TRUE)
str(lists)</pre>
```

makeNumeric

Change Character Variables to numeric

Description

Converts character variables, which contain only values, to numeric. Character variables containing letters are not converted. This avoids warnings, if conversion to numeric is attempted for variables, which contain characters.

Usage

makeNumeric(variable)

Arguments

variable Varia

Variable to be changed to numeric.

Value

Variable converted to numeric, if possible.

32 mergeData

Author(s)

Nicole Haag

Examples

```
a <- c("1", "2", "3", "4")
b <- c("1", "2", "x", "4")
makeNumeric(a)
makeNumeric(b)</pre>
```

mergeData

Merge Data Frames using one Key Variable

Description

Merges several data frames and matches them using one key variable

Usage

```
mergeData(newID = "ID", datList, oldIDs=NULL, addMbd = FALSE, writeLog=FALSE)
```

Arguments

| newID | character string containing the key variable's name in the merged dataset |
|----------|---|
| datList | list of data frames to be merged |
| oldIDs | character vector OR numeric vector containing either names of the key variables in datList or their column number in each dataframe in datList default is a vector containing replicates of the value of newID. |
| addMbd | logical; string "mbd" (missing by desgin) will be added instead of NA |
| writeLog | logical; if Logfile shall be written via sunk. |

Value

A data frame containing unique cases and unique variables. All cases and all variables that could be identified the original data frames will be kept and matched.

Author(s)

Karoline Sachse, Nicole Haag

Examples

```
data(inputDat)
str(inputDat)

mergedDataset <- mergeData("person-id", inputDat, c("idstud", "idstud", "idstud"), addMbd=TRUE)
str(mergedDataset)

mergedDataset <- mergeData("idstud", inputDat, writeLog=FALSE)
str(mergedDataset)</pre>
```

prepare.package 33

prepare.package

prepare.package

Description

prepares package

Usage

```
prepare.package ( source.folder = "p:/ZKD/development" ,
files ,
package.folder = "p:/ZKD/packages" ,
package.name ,
package.version )
```

Arguments

source.folder folder of R files

files character vector of R files that should be included in the package

package.version

version of package, must be in format "0.0.0"

Details

copies R files from source. folder to package folder copies "x.x.x" folder content to package folder modifies version and date in DESCRIPTION and automateModels creates ChangeLog

Author(s)

Martin Hecht

readDaemonXlsx

read xlsx-Files produced by ZKDaemon

Description

read xlsx-Files produced by ZKDaemon

Usage

```
readDaemonXlsx(filename)
```

Arguments

filename

A character string containing path, name and extension of .xlsx produced by ZKDaemon. Caution! Sheet order is important (see Datails).

34 recodeData

Details

Compulsory: 1st sheet: units. 2nd sheet: subunits. 3rd sheet: values. Optional: 4th sheet: unitRecodings. 5th sheet: savFiles. 6th sheet: newID. 7th sheet: aggregateMissings. 8th sheet: unitProperties. 9th sheet: property labels. 10th sheet: booklets.

Value

A list of data frames containing information that is required by automateDataPreparation

Author(s)

Karoline Sachse

Examples

str(inputList)

recodeData

Recode Datasets with Missing Values

Description

Recode datasets with special consideration of missing values.

Usage

```
recodeData(dat, values, subunits)
```

Arguments

dat A data frame

values A data frame with code information. See 'Details'. subunits A data frame with subunit information. See 'Details'.

Details

recodeData recodes data frames with special consideration of missing values. The results of recodeData will be written to a protocol file with sunk. recodeData will give warnings, if missing or incomplete recode informations are found. Values without recode information will NOT be recoded!

Examples of data frames values and subunits can be found via data(inputList)

Value

A data frame with recoded variables according to the specifications in values and subunits. Colnames will be the names specified in subunits\$subunitRecoded.

Author(s)

Martin Hecht, Christiane Penk, Nicole Haag

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References

http://code.google.com/p/zkdlib/wiki/MissingHandling

See Also

```
aggregateData, checkData
```

Examples

```
data(inputDat)
data(inputList)
# library(car)

dat1 <- inputDat[[1]] # get first dataset from inputDat
datRec <- recodeData(dat1, inputList$values, inputList$subunits)
str(datRec)</pre>
```

reinsort.col

reinsort.col

Description

insert columns of dataframe in specific position

Usage

```
reinsort.col ( dat , toreinsort , after )
```

Arguments

data data.frame on which operation should be performed

toreinsort column name(s) or numeric indicator(s) that should be relocated

after column name or numeric indicator after that toreinsort should be located

Value

data.frame

Author(s)

Martin Hecht

36 rmNAcols

rmNA

remove NA columns and rows from data

Description

remove columns and rows that are completely NA from data.frame or matrix

Usage

```
rmNA ( data , remove = TRUE , verbose = FALSE )
```

Arguments

data data.frame or matrix

remove if TRUE columns and rows are removed, if FALSE a list of identified columns and

rows is returned

verbose if TRUE removed columns and rows are printed on output window

Value

depends on option remove

Author(s)

Martin Hecht

See Also

```
rmNAcols, rmNArows
```

Examples

rmNAcols

remove NA columns from data

Description

remove columns that are completely or partially NA from data.frame or matrix

Usage

```
rmNAcols ( data , rows = NULL , tolerance = 0 , cumulate = TRUE , remove = TRUE , verbose = FALS
```

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Arguments

data data.frame or matrix

rows rows to include, can be a list of vectors to specify row subsets

tolerance number of non-NA cells that are "tolerated", can be a list corresponding to rows

cumulate if TRUE, tolerance is cumulated; if FALSE, exact tolerance is used

remove if TRUE, columns and rows are removed; if FALSE, identified columns are re-

turned

verbose if TRUE removed columns and rows are printed on output window

Value

depends on option remove

Author(s)

Martin Hecht

See Also

calls rmNA and rmNArows

Examples

```
# example matrix
# remove column with entirely NA (column 7)
rmNAcols( mat , verbose = TRUE )
\# remove column with NA on rows 3, 4, 5 (columns 5, 6, 7)
rmNAcols(mat, c(3,4,5), verbose = TRUE)
rmNAcols( mat , c(-1,-2,-6) , verbose = TRUE )
# tolerance=1 , 1 non-NA is permitted (columns 6 and 7)
rmNAcols( mat , tolerance=1 , verbose = TRUE )
# tolerance=6 , 6 non-NA are permitted (all columns are removed)
rmNAcols( mat , tolerance=6 , verbose = TRUE )
# do not cumulate / exact tolerance (column 1)
rmNAcols( mat , tolerance=6 , cumulate=FALSE , verbose = TRUE )
# two subsets of rows
rmNAcols( mat , rows = list( c(1, 2), c(4, 5) ) , verbose = TRUE )
# two subsets of rows with different tolerance
 rmNAcols( \ mat \ , \ rows = list( \ c(1), \ c(2, \ 3, \ 4, \ 5) \ ) \ , \ tolerance = list( \ 0 \ , \ 1 \ ) \ , \ verbose = TRUE \ ) 
# identify cols, no deletion
rmNAcols(\ mat\ ,\ rows\ =\ list(\ c(1,\ 2),\ c(3,\ 4,\ 5)\ )\ ,\ tolerance\ =\ list(\ 0\ ,\ 1\ )\ ,\ remove\ =\ FALSE\ )
```

38 rmNArows

| remove NA rows from data | Arows |
|---------------------------|--------|
| remove 141 rows from adia | ti OWS |

Description

remove rows that are completely or partially NA from data.frame or matrix

Usage

```
rmNArows ( data , cols = NULL , tolerance = 0 , cumulate = TRUE , remove = TRUE , verbose = FALS
```

Arguments

| data | data.frame or matrix |
|-----------|--|
| cols | columns to include, can be a list of vectors to specify column subsets |
| tolerance | number of non-NA cells that are "tolerated", can be a list corresponding to cols |
| cumulate | if TRUE, tolerance is cumulated; if FALSE, exact tolerance is used |
| remove | if TRUE, columns and rows are removed; if FALSE, identified rows are returned |
| verbose | if TRUE removed columns and rows are printed on output window |

Value

depends on option remove

Author(s)

Martin Hecht

See Also

calls rmNA and rmNAcols

Examples

rmNArows(mat , tolerance=5 , cumulate=FALSE , verbose = TRUE)

science1 39

```
rmNArows( mat , tolerance=5 , cumulate=FALSE , remove = FALSE )

# two subsets of columns
rmNArows( mat , cols = list( c(1, 2), c(4, 5) ) , verbose = TRUE )

# two subsets of columns with different tolerance
rmNArows( mat , cols = list( c(1), c(2, 3, 4, 5) ) , tolerance = list( 0 , 1 ) , verbose = TRUE )

# identify rows, no deletion
rmNArows( mat , cols = list( c(1), c(2, 3, 4, 5) ) , tolerance = list( 0 , 1 ) , remove = FALSE )
```

science1

Science achievement test data

Description

This data set contains responses of 420 students on 185 science items. Additional variables are included: id, grade, sex, booklet, track, version, and four dummy coded variables that indicate Track x Version groups. An incomplete block design was used with 4 booklets. Codes on items are: "0" - wrong "1" - right "mbd" - missing by design "mbi" - missing by intention "mir" - missing due to irregular response

Usage

```
data(science1)
```

Format

'data.frame': 420 obs. of 195 variables

Source

Simulated data

science1.context.vars Science achievement test data - Context variable names

Description

This vector contains the names of context variables in data set science1

Format

chr [1:9]

40 set.col.type

| | • . |
|----------|-----------|
| science1 | 1 † A M S |
| | |

Science achievement test data - Item names

Description

This vector contains the names items in data set science1

Format

```
chr [1:185]
```

science1.scales

Science achievement test data - Scale definition

Description

This data frame contains scale definitions for usage with automateModels and data set science1

Format

'data.frame': 185 obs. of 7 variables

set.col.type

set type of variable in data.frame

Description

```
converts type of column(s) to "character", "numeric", "logical", "integer" or "factor"
```

Usage

```
set.col.type ( data , col.type = list ( "character" = NULL ) , verbose = FALSE , ... )
```

Arguments

| data | data.frame |
|----------|--------------|
| cal type | nomed list o |

col.type named list of variable names that are to be converted. names of list is conversion type ("character" , "numeric" , "numeric.if.possible" , "logical" , "integer" or

"factor")

verbose if TRUE variables that have been converted are printed arguments to be passed to asNumericIfPossible

Details

use col.type="numeric.if.possible" if conversion to numeric should be tested upfront, see asNumericIfPossible for details

sortDfrByNames 41

Author(s)

Martin Hecht

Examples

```
str ( d <- data.frame ( "var1" = 1 , "var2" = TRUE , "var3" = FALSE , "var4" = as.factor ( 1 ) , "var5" = s
str ( set.col.type ( d ) )
str ( set.col.type ( d , list ( "numeric" = NULL ) ) )
str ( set.col.type ( d , list ( "character" = c ( "var1" , "var2" ) , "numeric" = "var3" , "logical" = "var3" t ( set.col.type ( d , list ( "numeric.if.possible" = NULL ) ) )
str ( set.col.type ( d , list ( "numeric.if.possible" = NULL ) , transform.factors = TRUE ) )
str ( set.col.type ( d , list ( "numeric.if.possible" = NULL ) , transform.factors = TRUE , maintain.factor</pre>
```

sortDfrByNames

sort data.frame by colnames and/or rownames

Description

specify new colnames and/or rownames order, data.frame is sorted in accordance

Usage

```
sortDfrByNames ( dfr , col.order = NULL , row.order = NULL , warn = TRUE )
```

Arguments

dfr data.frame

col.order character vector of colnames in new order row.order character vector of rownames in new order

warn logical, if TRUE warnings are printed on output window if col.order/row.order

do not correspond to colnames/rownames resp.

Value

data.frame

Author(s)

Martin Hecht

Examples

```
dfr <- data.frame ( matrix ( rnorm ( 100 ) , ncol = 10 ) )
colnames ( dfr ) <- paste ( "X" , 10:1 , sep = "" )
rownames ( dfr ) <- paste ( "X" , 11:2 , sep = "" )
dfr

# sort data.frame by 'col.order' and 'row.order'
sortDfrByNames ( dfr , paste ( "X" , 1:10 , sep = "" ) , paste ( "X" , 2:11 , sep = "" ) )</pre>
```

42 source.it.all2

source.it.all source.it.all

Description

sources *.R files of folder

Usage

```
source.it.all ( folder="p:/ZKD/development" , develop.modules = NULL , return.stable = FALSE )
```

Arguments

folder folder with *.R files

develop.modules

character vector of R files that should be sourced in development status return.stable if TRUE nothing is sourced and a vector of all stable versions is returned

Value

return.stable = FALSE

sources R files

return.stable = TRUE

character vector of stable R files

Author(s)

Christiane Penk, Martin Hecht

source.it.all2 source.it.all2

Description

sources *.R files of folder

Usage

```
source.it.all2 ( folder="p:/ZKD/development" , use.zkd.conv = TRUE , development = TRUE , development
```

Arguments

folder folder with *.R files

development if TRUE development versions are sourced (if non-existent the latest stable is

sourced or nothing is sourced, see option development.only\ if FALSE stable

versions are sourced

use.zkd.conv if TRUE R files in folder are checked to be consisten with specific ("zkd") ver-

sioning convention \ if FALSE all R files in folder are sourced

development.only

if TRUE only development versions are sourced \ if FALSE stable versions are

included

exclude character vector of R files that should not be sourced

sunk 43

Value

sources R files

Author(s)

Martin Hecht, Christiane Penk

sunk sunk

Description

writes output to file

Usage

```
sunk (cmd = NULL, path = NULL, write = TRUE, console.output = TRUE, new.file = FALSE, text
```

Arguments

cmd character string of element to write, may be either text (e.g. "write me to file")

or a function call (e.g. "summary(lm)")

path (folder and name) to output file if NULL path is defaulted to getwd()+"sunk.txt"

all environments are searched for sunk.path, if sunk.path is found (exists), it is

used

write logical, if TRUE (default) output is written to file

console.output logical, if TRUE (default) output is displayed on console

new.file logical, if TRUE the output file is created if FALSE (default) output is appended

to existing file

text.on.error logical, sunk checks if the character string 'cmd' is an evaluatable expression

if TRUE (default), 'cmd' is treated as text if an error occurs when trying to

evaluate string if FALSE, sunk stops on errors/not evaluatable expressions

text.out.method

choose "cat" (default) or "print" as the output method for text

Value

writes to disk

Author(s)

Martin Hecht

44 writeSpss

| writeSpss | Export Datasets to SPSS |
|-----------|-------------------------|
| | |

Description

Writes data and SPSS syntax files.

Usage

Arguments

| dat | A data frame |
|--------------|--|
| values | A data frame with code information. See 'Details'. |
| subunits | A data frame with subunit information. See 'Details'. |
| units | A data frame with unit information. See 'Details'. |
| filedat | A character string with the name of the output data file. |
| filesps | A character string with the name of the output syntax file. |
| missing.rule | A list containing recode information for character missings. See 'References' for description of default values. |
| path | A character string containing the path of the output file. The value in path is appended to filedat and filesps. By default, files are written to the current R working directory. If path=NULL then no file path appending is done. |
| sep | The separator between the data fields. |
| dec | The decimal separator for numerical data. |
| silent | A logical flag stating whether the names of the files should be printed. |

Details

This function automates most of the work needed to export a dataset to SPSS. It uses a modified version of writeForeignSPSS() from the foreign() package and of mids2spss() from the mice package. The modified version allows for a choice of the field and decimal separators, makes some improvements to the formatting and provides variable labels and value labels according to the information in the data frames values, subunits and units.

Examples of data frames values, subunits and units can be found on data(inputList)

The SPSS syntax file has the proper file names and separators set, so in principle it should run and read the data without alteration. SPSS is more strict than R with respect to the paths. Always use the full path, otherwise SPSS may not be able to find the data file.

Value

Used for its side effects. The return value is NULL.

yen.q3 45

Author(s)

Nicole Haag

References

http://code.google.com/p/zkdlib/wiki/MissingHandling

|--|--|

Description

Q3 statistics

Usage

```
yen.q3 ( dat , theta , b , progress = T )
```

Arguments

```
dat bla theta bla b bla progress bla
```

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