Package 'mrdwabmisc'

February 12, 2013

Title Miscellaneous R functions, mostly for data processing

Type Package

Index

 stringseed.sampling
 16

 table2df
 18

21

2 mrdwabmisc-package

mrdwabmisc-package mrdwabmisc

Description

Miscellaneous R functions, some utility, and others to clean and organize data.

Details

Package: mrdwabmisc
Type: Package
Version: 1.0
Date: 2013-01-22
License: GPL-2

Author(s)

Ananda Mahto

Maintainer: Ananda Mahto <mrdwab@gmail.com>

See Also

AMsnippets contains a related set of utility functions, but the packages do not depend on each other.

concat.split 3

concat.split	Split concatenated cells in a data.frame	
--------------	--	--

Description

The concat.split function takes a column with multiple values, splits the values into a list or into separate columns, and returns a new data.frame.

Usage

```
concat.split(data, split.col, sep = ",",
   structure = "compact", mode = NULL, drop.col = FALSE,
   fixed = FALSE)
```

Arguments

data	The source data.frame
split.col	The variable that needs to be split; can be specified either by the column number or the variable name.
sep	The character separating each value (defaults to ",").
structure	Can be either "compact", "expanded", or "list". Defaults to "compact". See Details.
mode	Can be either binary or value (where binary is default and it recodes values to 1 or NA, like Boolean data, but without assuming 0 when data is not available). This setting only applies when structure = "expanded"; an warning message will be issued if used with other structures.
drop.col	Logical (whether to remove the original variable from the output or not). Defaults to TRUE.
fixed	Is the input for the sep value fixed, or a regular expression? See Details.

Details

structure

- "compact" creates as many columns as the maximum length of the resulting split. This is the most useful general-case application of this function.
- When the input is numeric, "expanded" creates as many columns as the maximum value of the input data. This is most useful when converting to mode = "binary".
- "list" creates a single new column that is structurally a list within a data. frame.

fixed When structure = "expanded" or structure = "list", it is possible to supply a a regular expression containing the characters to split on. For example, to split on ",", ";", or "|", you can set sep = ", |; | | |" or sep = "[,; |]", and fixed = FALSE to split on any of those characters.

4 concat.split

Note

If using structure = "compact", the value for sep can only be a single character. See the "Advanced Usage" example of how to specify multiple characters for batch conversion of columns.

Author(s)

Ananda Mahto

References

- See http://stackoverflow.com/q/10100887/1270695
- The "condensed" setting was inspired by an answer from David Winsemius to a question at Stack Overflow. See: http://stackoverflow.com/a/13924245/1270695

```
## Load some data
data(concatenated)
head(concat.test)
# Split up the second column, selecting by column number
head(concat.split(concat.test, 2))
# ... or by name, and drop the offensive first column
head(concat.split(concat.test, "Likes", drop.col = TRUE))
# The "Hates" column uses a different separator
head(concat.split(concat.test, "Hates", sep = ";", drop.col = TRUE))
# You'll get a warning here, when trying to retain the original values
head(concat.split(concat.test, 2, mode = "value", drop.col = TRUE))
# Try again. Notice the differing number of resulting columns
head(concat.split(concat.test, 2, structure = "expanded",
     mode = "value", drop.col = TRUE))
# Let's try splitting some strings... Same syntax
head(concat.split(concat.test, 3, drop.col = TRUE))
# Split up the "Likes column" into a list variable; retain original column
head(concat.split(concat.test, 2, structure = "list", drop.col=FALSE))
# View the structure of the output for the first 10 rows to verify
# that the new column is a list; note the difference between "Likes"
# and "Likes_list".
str(concat.split(concat.test, 2, structure = "list",
  drop.col=FALSE)[1:10, c(2, 5)])
# ADVANCED USAGE ###
# Show just the first few lines, compact structure
```

df.sorter 5

df.sorter

Sort a data. frame by rows or columns

Description

The df. sorter function allows you to sort a data. frame by columns or rows or both. You can also quickly subset data columns by using the var. order argument.

Usage

```
df.sorter(data, var.order = names(data), col.sort = NULL,
    at.start = TRUE)
```

Arguments

data The source data.frame.

var.order The new order in which you want the variables to appear. See Details

col.sort The columns within which there is data that need to be sorted. See Details.

Should the pattern matching be from the start of the variable name? Defaults to TRUE.

Details

var.order

- Defaults to names (data), which keeps the variables in the original order.
- Variables can be referred to either by a vector of their index numbers or by a vector of the variable name; partial name matching also works, but requires that the partial match identifies similar columns uniquely (see Examples). Basic subsetting can also be done using var.order simply by omitting the variables you want to drop.

col.sort

• Defaults to NULL, which means no sorting takes place. Variables can be referred to either by a vector of their index numbers or by a vector of the variable names; full names must be provided.

6 multi.freq.table

Note

If you are sorting both by variables and within the columns and using numeric indexes as opposed to variable names, the col.sort order should be based on the location of the columns in the new data.frame, not the original data.frame.

Author(s)

Ananda Mahto

Examples

```
# Make up some data
set.seed(1)
dat = data.frame(id = rep(1:5, each=3), times = rep(1:3, 5),
                 measure1 = rnorm(15), score1 = sample(300, 15),
                 code1 = replicate(15, paste(sample(LETTERS[1:5], 3),
                                             sep="", collapse="")),
                 measure2 = rnorm(15), score2 = sample(150:300, 15),
                 code2 = replicate(15, paste(sample(LETTERS[1:5], 3),
                                             sep="", collapse="")))
# Preview your data
dat
# Change the variable order, grouping related columns
# Note that you do not need to specify full variable names,
     just enough that the variables can be uniquely identified
head(df.sorter(dat, var.order = c("id", "ti", "cod", "mea", "sco")))
# As above, but sorted by 'times' and then 'id'
head(df.sorter(dat,
               var.order = c("id", "tim", "cod", "mea", "sco"),
               col.sort = c(2, 1))
# Drop 'measure1' and 'measure2', sort by 'times', and 'score1'
head(df.sorter(dat,
               var.order = c("id", "tim", "sco", "cod"),
               col.sort = c(2, 3))
# Just sort by columns, first by 'times' then by 'id'
head(df.sorter(dat, col.sort = c("times", "id")))
# Pattern matching anywhere in the variable name
head(df.sorter(dat, var.order= "co", at.start=FALSE))
```

multi.freq.table Ta

 $\it Tabulates\ columns\ from\ a\ data.$ frame $\it containing\ multiple-response\ data$

multi.freq.table 7

Description

The multi.freq.table function takes a data.frame containing Boolean responses to multiple response questions and tabulates the number of responses by the possible combinations of answers.

Usage

```
multi.freq.table(data, sep = "", boolean = TRUE,
factors = NULL, NAto0 = TRUE, basic = FALSE,
dropzero = TRUE, clean = TRUE)
```

Arguments

data	The multiple responses that need to be tabulated.
sep	The desired separator for collapsing the combinations of options; defaults to "" (collapsing with no space between each option name).
boolean	Are you tabulating boolean data (see dat Examples)? Defaults to TRUE.
factors	If you are trying to tabulate non-boolean data, and the data are not factors, you can specify the factors here (see dat2 Examples). Defaults to NULL and is not used when boolean = TRUE.
NAto0	Should NA values be converted to 0? Defaults to TRUE, in which case, the number of valid cases should be the same as the number of cases overall. If set to FALSE, any rows with NA values will be dropped as invalid cases. Only applies when boolean = TRUE.
basic	Should a basic table of each item, rather than combinations of items, be created? Defaults to FALSE.
dropzero	Should combinations with a frequency of zero be dropped from the final table? Defaults to TRUE. Does not apply when boolean = TRUE.
clean	Should the original tabulated data be retained or dropped from the final table? Defaults to TRUE (drop). Does not apply when boolean = TRUE.

Details

In addition to tabulating the *frequency* (Freq), there are two other columns in the output: *Percent of Responses* (Pct.of.Resp) and *Percent of Cases* (Pct.of.Cases).

Percent of Responses is the frequency divided by the total number of answers provided; this column should sum to 100 table is generated and there are cases where a respondent did not select any option, the Percent of Responses value would be more than 100 frequency divided by the total number of valid cases; this column would most likely sum to more than 100 a basic table is produced since each respondent (case) can select multiple answers, but should sum to 100 other tables.

Author(s)

Ananda Mahto

References

apply shortcut for creating the Combn column in the output by Justin. See: http://stackoverflow.com/q/11348391/1270695 and http://stackoverflow.com/q/11622660/1270695

8 multi.freq.table

```
## ======== ##
## ====== BOOLEAN DATA ======= ##
# Make up some data
set.seed(1)
dat <- data.frame(A = sample(c(0, 1), 20, replace=TRUE),
                B = sample(c(0, 1, NA), 20,
                          prob=c(.3, .6, .1), replace=TRUE),
                C = sample(c(0, 1, NA), 20,
                          prob=c(.7, .2, .1), replace=TRUE),
                D = sample(c(0, 1, NA), 20,
                          prob=c(.3, .6, .1), replace=TRUE),
                E = sample(c(0, 1, NA), 20,
                          prob=c(.4, .4, .2), replace=TRUE))
# View your data
dat
# How many cases have "NA" values?
table(is.na(rowSums(dat)))
# Apply the function with all defaults accepted
multi.freq.table(dat)
# Tabulate only on variables "A", "B", and "D", with a different
# separator, keep any zero frequency values, and keeping the
# original tabulations. There are no solitary "D" responses.
multi.freq.table(dat[c(1, 2, 4)], sep="-", dropzero=FALSE, clean=FALSE)
# As above, but without converting "NA" to "0".
# Note the difference in the number of valid cases.
multi.freq.table(dat[c(1, 2, 4)], NAto0=FALSE,
                sep="-", dropzero=FALSE, clean=FALSE)
# View a basic table.
multi.freq.table(dat, basic=TRUE)
## ========= ##
## ======= NON-BOOLEAN DATA ======== ##
# Make up some data
dat2 <- structure(list(Reason.1 = c("one", "one", "two", "one", "two",</pre>
                                 "three", "one", "one", NA, "two"),
                     Reason.2 = c("two", "three", "three", NA, NA,
                                 "two", "three", "two", NA, NA),
                    Reason.3 = c("three", NA, NA, NA, NA,
                                 NA, NA, "three", NA, NA)),
                     .Names = c("Reason.1", "Reason.2", "Reason.3"),
                     class = "data.frame",
                     row.names = c(NA, -10L))
```

RandomNames 9

RandomNames

Generate random names

Description

The RandomNames function uses data from the *Genealogy Data: Frequently Occurring Surnames from Census 1990–Names Files* web page to generate a data. frame with random names.

Usage

```
RandomNames(N = 100, cat = NULL, gender = NULL,
   MFprob = NULL, dataset = NULL)
```

Arguments

N The number of random names you want. Defaults to 100.

Do you want "common" names, "rare" names, names with an "average" frequency, or some combination of these? Should be specified as a character vec-

tor (for example, c("rare", "common")). Defaults to NULL, in which case all

names are used as the sample frame.

gender Do you want first names from the "male" dataset, the "female" dataset, or from

all available names? Should be specified as a quoted string (for example, "male"). Defaults to NULL, in which case all available first names are used as the sample

frame.

MFprob What proportion of the sample should be male names and what proportion should be female? Specify as a numeric vector that sums to 1 (for example,

should be female? Specify as a numeric vector that sums to 1 (for example, c(.6, .4)). The first number represents the probability of sampling a "male" first name, and the second number represents the probability of sampling a "female" name. This argument is not used if only one gender has been specified in the previous argument. Defaults to NULL, in which case, the probability used is

c(.5, .5).

10 RandomNames

dataset

What do you want to use as the dataset of names from which to sample? A default dataset is provided that can generate over 400 million unique names. See the "Dataset Details" note for more information.

Note

Dataset Details This function samples from a provided dataset of names. By default, it uses the data from the Genealogy Data: Frequently Occurring Surnames from Census 1990–Names Files web page. Those data have been converted to list named "CensusNames1990" containing three data.frames (named "surnames", "malenames", and "femalenames").

Alternatively, you may provide your own data in a list formatted according to the following specifications (see the "myCustomNames" data in the "Examples*" section). *Please remember that R is case sensitive!*

- This must be a named list with three items: "surnames", "malenames", and "femalenames".
- The contents of each list item is a data.frame with at least the following named columns: "Name" and "Category".
- Acceptable values for "Category" are "common", "rare", and "average".

Author(s)

Ananda Mahto

References

- See http://www.census.gov/genealogy/www/data/1990surnames/names_files.html for source of data
- Inspired by the online Random Name Generator http://random-name-generator.info/

row.extractor 11

```
myCustomNames <- list(
  surnames = data.frame(
    Name = LETTERS[1:26],
    Category = c(rep("rare", 10), rep("average", 10), rep("common", 6))),
  malenames = data.frame(
    Name = letters[1:10],
    Category = c(rep("rare", 4), rep("average", 4), rep("common", 2))),
  femalenames = data.frame(
    Name = letters[11:26],
    Category = c(rep("rare", 8), rep("average", 4), rep("common", 4))))
  str(myCustomNames)</pre>
RandomNames(N = 15, dataset = myCustomNames)
```

row.extractor

Extract min/median/max/quantile rows from a data.frame

Description

The row.extractor function takes a data. frame and extracts rows with the min, median, or max values of a given variable, or extracts rows with specific quantiles of a given variable.

Usage

```
row.extractor(data, extract.by, what = "all")
```

Arguments

data The source data. frame.

extract.by The column which will be used as the reference for extraction; can be specified

either by the column number or the variable name.

what Options are "min" (for all rows matching the minimum value), "median" (for

the median row or rows), "max" (for all rows matching the maximum value), or "all" (for min, median, and max); alternatively, a numeric vector can be specified with the desired quantiles, for instance c(0, .25, .5, .75, 1).

Author(s)

Ananda Mahto

References

- which.quantile function by cheleites: http://stackoverflow.com/users/755257/cheleites
- See: http://stackoverflow.com/q/10256503/1270695

12 sample.size

See Also

```
min, max, median, which.min, which.max, quantile
```

Examples

```
# Make up some data
set.seed(1)
dat = data.frame(V1 = 1:50, V2 = rnorm(50),
                V3 = round(abs(rnorm(50)), digits=2),
                V4 = sample(1:30, 50, replace=TRUE))
# Get a sumary of the data
summary(dat)
# Get the rows corresponding to the 'min', 'median', and 'max' of 'V4'
row.extractor(dat, 4)
# Get the 'min' rows only, referenced by the variable name
row.extractor(dat, "V4", "min")
# Get the 'median' rows only. Notice that there are two rows
     since we have an even number of cases and true median
     is the mean of the two central sorted values
row.extractor(dat, "V4", "median")
# Get the rows corresponding to the deciles of 'V3'
row.extractor(dat, "V3", seq(0.1, 1, 0.1))
```

sample.size

Determine the optimal sample size of a given population

Description

The sample.size function either calculates the optimum survey sample size when provided with a population size, or the confidence interval of using a certain sample size with a given population. It can be used to generate tables (data.frames) of different combinations of inputs of the following arguments, which can be useful for showing the effect of each of these in sample size calculation.

Usage

```
sample.size(population, samp.size = NULL, c.lev = 95,
    c.int = NULL, what = "sample", distribution = 50)
```

Arguments

population The population size for which a sample size needs to be calculated.

samp.size The sample size. This argument is only used when calculating the confidence interval, and defaults to NULL.

sample.size 13

c.lev	The desired confidence level. Defaults to a reasonable 95%.
c.int	The confidence interval. This argument is only used when calculating the sample size. If not specified when calculating the sample size, defaults to 5% and a message is provided indicating this; this is also the default action if c.int = NULL.
what	Should the function calculate the desired sample size or the confidence interval? Accepted values are "sample" and "confidence" (quoted), and defaults to "sample".
distribution	Response distribution. Defaults to 50% (distribution = 50), which will give you the largest sample size.

Note

From a teaching perspective, the function can be used to easily make tables which demonstrate how the sample size or confidence interval change when different inputs change. See the "Advanced Usage" examples. The following formulae were used in this function:

$$ss = \frac{-Z^2 \times p \times (1-p)}{c^2}$$

$$pss = \frac{ss}{1 + \frac{ss - 1}{pop}}$$

Author(s)

Ananda Mahto

References

- See the 2657 Productions News site for how this function progressively developed: http://news.mrdwab.com/2010/09/10/a-sample-size-calculator-function-for-r/
- The sample.size function is based on the following formulas from the Creative Research Systems web page *Sample size formulas for our sample size calculator*: http://www.webcitation.org/69kNjMuKe

```
# What should our sample size be for a population of 300?
# All defaults accepted.
sample.size(population = 300)

# What sample should we take for a population of 300
# at a confidence level of 97%?
sample.size(population = 300, c.lev = 97)

# What about if we change our confidence interval?
sample.size(population = 300, c.int = 2.5, what = "sample")
# What about if we want to determine the confidence interval
```

14 stratified

```
of a sample of 140 from a population of 300? A confidence
   level of 95% is assumed.
sample.size(population = 300, samp.size = 140, what = "confidence")
## ========= ##
## ====== ADVANCED USAGE ======= ##
# What should the sample be for populations of 300 to 500 by 50?
sample.size(population=c(300, 350, 400, 450, 500))
# How does varying confidence levels or confidence intervals
   affect the sample size?
sample.size(population=300,
           c.lev=rep(c(95, 96, 97, 98, 99), times = 3),
           c.int=rep(c(2.5, 5, 10), each=5))
# What is are the confidence intervals for a sample of
   150, 160, and 170 from a population of 300?
sample.size(population=300,
           samp.size = c(150, 160, 170),
           what = "confidence")
```

stratified

 $Sample\ from\ a\ {\it data.frame}\ according\ to\ a\ stratification\ variable$

Description

The stratified function samples from a data.frame in which one of the columns can be used as a "stratification" or "grouping" variable. The result is a new data.frame with the specified number of samples from each group.

Usage

```
stratified(df, group, size, seed = NULL, ...)
```

Arguments

df

The source data.frame.

group

Your grouping variables. Generally, if you are using more than one variable to create your "strata", you should list them in the order of *slowest* varying to *quickest* varying. This can be a vector of names or column indexes.

size

The desired sample size.

- If size is a value between 0 and 1 expressed as a decimal, size is set to be proportional to the number of observations per group.
- If size is a single positive integer, it will be assumed that you want the same number of samples from each group.

stratified 15

If size is a vector, the function will check to see whether the length of the
vector matches the number of groups and use those specified values as the
desired sample sizes. The values in the vector should be in the same order
as you would get if you tabulated the grouping variable (usually alphabetic order); alternatively, you can name each value to ensure it is properly
matched.

seed

The seed that you want to use (using set. seed within the function, if any. Defaults to NULL.

. . .

Further arguments to be passed to the sample function.

Note

Slightly different sizes than requested

Because of how computers deal with floating-point arithmetic, and because R uses a "round to even" approach, the size per strata that results when specifying a proportionate sample may be slightly higher or lower per strata than you might have expected.

"Seed" argument

This is different from using set.seed before using the function. Setting a seed using this argument is equivalent to using set.seed each time that you go to take a sample from a different group (in other words, the same seed is used for each group).

The inclusion of a seed argument is mostly a matter of convenience, to be able to have a single seed with which the samples can be verified later. However, by using the seed argument, the same seed is used to sample from each group. This may be a problem if there are many groups that have the same number of observations, since it means that the same observation number will be selected from each of those grops. For instance, if group "AA" and "DD" both had the same number of observations (say, 5) and you were sampling 3 cases using a seed of 1, the second, fifth, and fourth observation would be taken from each of those groups. To avoid this, you can set the seed using set.seed *before* you run the stratified function.

As a user, you need to weigh the benefits and drawbacks of setting the seed *before* running the function as opposed to setting the seed *with* the function. Setting the seed before would be useful if there are several groups with the same number of observations; however, in the slim chance that you need to verify the samples manually, you *may* run into problems.

Author(s)

Ananda Mahto

References

The evolution of this function can be traced at the following links. The version in this package is entirely reworked and does not require an additional package to be loaded.

- http://news.mrdwab.com/2011/05/15/stratified-random-sampling-in-r-beta/
- http://news.mrdwab.com/2011/05/20/stratified-random-sampling-in-r-from-a-data-frame/
- http://stackoverflow.com/a/9714207/1270695

16 stringseed.sampling

See Also

strata

```
# Generate a couple of sample data.frames to play with
set.seed(1)
dat1 \leftarrow data.frame(ID = 1:100,
                   A = sample(c("AA", "BB", "CC", "DD", "EE"), 100, replace = TRUE),
                   B = rnorm(100), C = abs(round(rnorm(100), digits=1)),
                   D = sample(c("CA", "NY", "TX"), 100, replace = TRUE),
                   E = sample(c("M", "F"), 100, replace = TRUE))
dat2 <- data.frame(ID = 1:20,</pre>
                   A = c(rep("AA", 5), rep("BB", 10),
                         rep("CC", 3), rep("DD", 2)))
# What do the data look like in general?
summary(dat1)
summary(dat2)
# Let's take a 10% sample from all -A- groups in dat1, seed = 1
stratified(dat1, "A", .1, seed = 1)
# Let's take 5 samples from all -D- groups in dat1,
# seed = 1, specified by column number
stratified(dat1, group = 5, size = 5, seed = 1)
# Let's take a sample from all -A- groups in dat1, seed = 1,
   where we specify the number wanted from each group
stratified(dat1, "A", size = c(3, 5, 4, 5, 2), seed = 1)
# Use a two-column strata: -E- and -D-
# -E- varies more slowly, so it is better to put that first
stratified(dat1, c("E", "D"), size = .15, seed = 1)
# Use a three-column strata: -E-, -D-, and -A-
s.out <- stratified(dat1, c("E", "D", "A"), size = 2, seed = 1)</pre>
list(head(s.out), tail(s.out))
# How many samples were taken from each strata?
table(interaction(s.out[c("E", "D", "A")]))
# Can we verify the message about group sizes?
names(which(table(interaction(dat1[c("E", "D", "A")])) < 2))</pre>
names(which(table(interaction(s.out[c("E", "D", "A")])) < 2))</pre>
```

stringseed.sampling 17

Description

The stringseed.sampling function is designed as a batch sampling function that allows the user to specify any alphanumeric input as the seed *per sample in the batch*.

Usage

```
stringseed.sampling(seedbase, N, n, write.output = FALSE)
```

Arguments

seedbase A vector of seeds to be used for sampling.

N The "population" from which to draw the sample.

n The desired number of samples.

write.output Logical. Should the output be written to a file? Defaults to FALSE. If TRUE,

a csv file is written with the sample "metadata", and a plain text file is written with the details of the resulting sample. The names of the files written are "Sample frame generated on {date the script was run} .csv" and "Samples generated on {date the script was run} .txt" and will be

found in your current working directory.

Value

This function returns a list with the class "stringSeedSampling" and uses a custom print method. The list items are:

- input: The "metadata" to remind you of your input parameters.
- samples: The samples resulting from the specified input parameters. In the case of batch sampling being used, samples will be a named list.

Use print.default(your-object-here) to view the underlying list.

Author(s)

Ananda Mahto

References

See: http://stackoverflow.com/q/10910698/1270695

See Also

digest

18 table2df

Examples

```
# We'll use a data.frame with a list of village names, the population,
   and the desired samples as our columns. The function will use the
   village names to generate a unique seed for each village before
  drawing the sample.
myListOfPlaces <- data.frame(</pre>
 villageName = c("Melakkal", "Sholavandan", "T. Malaipatti"),
 population = c(120, 130, 140),
  requiredSample = c(30, 25, 12))
myListOfPlaces
stringseed.sampling(seedbase = myListOfPlaces$villageName,
                    N = myListOfPlaces$population,
                    n = myListOfPlaces$requiredSample)
# Manual verification of the samples generated for Melakkal village
    (for which the automatically generated seed was 1331891848)
set.seed(1331891848)
sample(120, 30)
# What about using the function on a single input?
stringseed.sampling("Santa Barbara", 1920, 100)
```

table2df

Convert table objects to data.frames

Description

The table2df function takes an object of "class" table, ftable, and xtabs and converts them to data.frames, while retaining as many of the name details as possible.

Usage

```
table2df(mytable, as.multitable = FALSE,
  direction = "wide")
```

Arguments

mytable

The table object you want to convert into a data. frame. This can be an object in your workspace, or you can make the call to table, ftable, or xtabs as the mytable argument to this function.

as.multitable

Logical; defaults to FALSE. Some methods, for instance xtabs and table, will create an array of tables as the output when more than two variables are being tabulated.

- If as.multitable is TRUE, the function will return a list of data.frames.
- If as.multitable is FALSE, the function will convert the object to an ftable object before performing the transformation.

table2df

direction

Can be either "long" or "wide".

• If "long", the frequencies will all be tabulated into a single column. This is the same behavior you will generally get if you used as .data.frame on a table object.

• If "wide", the tabular format is retained.

Author(s)

Ananda Mahto

References

The expand.grid method for remaking the columns from an ftable was described by Kohske at http://stackoverflow.com/a/6463137/1270695.

See Also

```
table, ftable, xtabs
```

```
# Make up some data:
set.seed(1)
handedness <- data.frame(</pre>
  gender = sample(c("Female", "Male", "Unknown"), 200, replace = TRUE),
  handedness = sample(c("Right", "Left", "Ambidextrous"),
                      200, replace = TRUE, prob = c(.7, .2, .1)),
  fav.col = sample(c("Red", "Orange", "Yellow", "Green", "Blue",
                    "Indigo", "Violet", "Black", "White"),
                   200, replace = TRUE),
  fav.shape = sample(c("Triangle", "Circle", "Square", "Pentagon", "Hexagon",
                      "Oval", "Octagon", "Rhombus", "Trapezoid"),
                      200, replace = TRUE),
  computer = sample(c("Win", "Mac", "Lin"), 200, replace = TRUE,
                    prob = c(.5, .25, .25))
# Preview the data
list(head(handedness), tail(handedness))
# A very basic table
HT1 <- with(handedness, table(gender, handedness))</pre>
table2df(HT1)
#'table2df(HT1, direction = "long")
# Another basic table
HT2 <- with(handedness, table(fav.col, computer))</pre>
HT2
table2df(HT2)
# This will create multiple tables, one for each possible computer value
HT3 <- with(handedness, table(gender, fav.col, computer))</pre>
```

20 table2df

```
HT3
# Default settings
table2df(HT3)
# As a list of data.frames
table2df(HT3, as.multitable = TRUE)
# As above, but with the output in long format
# Only showing the first three lines of each data.frame
lapply(table2df(HT3, as.multitable = TRUE, direction = "long"), head, 3)
# Applied to an ftable
HT4 <- ftable(handedness,
             col.vars="fav.col",
             row.vars=c("gender", "computer"))
HT4
table2df(HT4)
## ======== ##
## ======= OTHER EXAMPLES ======= ##
## Not run:
table2df(xtabs(cbind(ncases, ncontrols) ~ ., data = esoph))
table2df(xtabs(cbind(ncases, ncontrols) ~ ., data = esoph),
        direction = "long")
table2df(xtabs(cbind(ncases, ncontrols) \sim ., data = esoph),
        as.multitable = TRUE, direction = "long")
table2df(xtabs(cbind(ncases, ncontrols) \sim ., data = esoph),
        as.multitable = TRUE, direction = "wide")
## End(Not run)
```

Index

```
*Topic package
    mrdwabmisc-package, 2
{\bf AMsnippets, 2}\\
concat.split, 3
data.frame, 3, 5, 7, 9, 14, 18
df.sorter, 5, 5
digest, 17
ftable, 18, 19
list, 3
max, 12
median, 12
min, 12
mrdwabmisc (mrdwabmisc-package), 2
mrdwabmisc-package, 2
multi.freq.table, 6, 7
quantile, 12
RandomNames, 9, 9
row.extractor, 11, 11
sample, 15
sample.size, 12
set.seed, 15
strata, 16
stratified, 14, 14, 15
stringseed.sampling, 16, 17
table, 18, 19
table2df, 18, 18
which.max, 12
which.min, 12
xtabs, 18, 19
```